WO 01/23604



TITLE OF THE INVENTION

HIGHLY CONSERVED GENES AND THEIR USE TO GENERATE SPECIESSPECIFIC, GENUS-SPECIFIC, FAMILY-SPECIFIC, GROUP-SPECIFIC AND
UNIVERSAL NUCLEIC ACID PROBES AND AMPLIFICATION PRIMERS TO
RAPIDLY DETECT AND IDENTIFY ALGAL, ARCHAEAL, BACTERIAL,
FUNGAL AND PARASITICAL MICROORGANISMS FROM CLINICAL
SPECIMENS FOR DIAGNOSIS

BACKGROUND OF THE INVENTION

Classical methods for the identification of microorganisms

Microorganisms are classically identified by their ability to utilize different substrates as a source of carbon and nitrogen through the use of biochemical tests such as the API20ETM system (bioMérieux). For susceptibility testing, clinical microbiology laboratories use methods including disk diffusion, agar dilution and broth microdilution. Although identifications based on biochemical tests and antibacterial susceptibility tests are cost-effective, generally two days are required to obtain preliminary results due to the necessity of two successive overnight incubations to identify the bacteria from clinical specimens as well as to determine their susceptibility to antimicrobial agents. There are some commercially available automated systems (i.e. the MicroScanTM system from Dade Behring and the VitekTM system from bioMérieux) which use sophisticated and expensive apparatus for faster microbial identification and susceptibility testing (Stager and Davis, 1992, Clin. Microbiol. Rev. 5:302-327). These systems require shorter incubation periods, thereby allowing most bacterial identifications and susceptibility testing to be performed in less than 6 hours. Nevertheless, these

faster systems always require the primary isolation of the bacteria or fungi as a pure culture, a process which takes at least 18 hours for a pure culture or 2 days for a mixed culture. So, the shortest time from sample reception to identification of the pathogen is around 24 hours. Moreover, fungi other than yeasts are often difficult or very slow to grow from clinical specimens. Identification must rely on labor-intensive techniques such as direct microscopic examination of the specimens and by direct and/or indirect immunological assays. Cultivation of most parasites is impractical in the clinical laboratory. Hence, microscopic examination of the specimen, a few immunological tests and clinical symptoms are often the only methods used for an identification that frequently remains presumptive.

The fastest bacterial identification system, the autoSCAN-Walk-AwayTM system (Dade Behring) identifies both gram-negative and gram-positive bacterial species from standardized inoculum in as little as 2 hours and gives susceptibility patterns to most antibiotics in 5 to 6 hours. However, this system has a particularly high percentage (i.e. 3.3 to 40.5%) of non-conclusive identifications with bacterial species other than *Enterobacteriaceae* (Croizé J., 1995, Lett. Infectiol. 10:109-113; York *et al.*, 1992, J. Clin. Microbiol. 30:2903-2910). For *Enterobacteriaceae*, the percentage of non-conclusive identifications was 2.7 to 11.4%. The list of microorganisms identified by commercial systems based on classical identification methods is given in Table 15.

A wide variety of bacteria and fungi are routinely isolated and identified from clinical specimens in microbiology laboratories. Tables 1 and 2 give the incidence for the most commonly isolated bacterial and fungal pathogens from various types of clinical specimens. These pathogens are the main organisms associated with nosocomial and community-acquired human infections and are therefore considered the most clinically important.

Clinical specimens tested in clinical microbiology laboratories

Most clinical specimens received in clinical microbiology laboratories are urine and blood samples. At the microbiology laboratory of the Centre Hospitalier de l'Université Laval (CHUL), urine and blood account for approximately 55% and 30% of the specimens received, respectively (Table 3). The remaining 15% of clinical specimens comprise various biological fluids including sputum, pus, cerebrospinal fluid, synovial fluid, and others (Table 3). Infections of the urinary tract, the respiratory tract and the bloodstream are usually of bacterial etiology and require antimicrobial therapy. In fact, all clinical samples received in the clinical microbiology laboratory are tested routinely for the identification of bacteria and antibiotic susceptibility.

Conventional pathogen identification from clinical specimens

Urine specimens

The search for pathogens in urine specimens is so preponderant in the routine microbiology laboratory that a myriad of tests have been developed. However, the gold standard remains the classical semi-quantitative plate culture method in which 1 µL of urine is streaked on agar plates and incubated for 18-24 hours. Colonies are then counted to determine the total number of colony forming units (CFU) per liter of urine. A bacterial urinary tract infection (UTI) is normally associated with a bacterial count of 10⁷ CFU/L or more in urine. However, infections with less than 10⁷ CFU/L in urine are possible, particularly in patients with a high incidence of diseases or those catheterized (Stark and Maki, 1984, N. Engl. J. Med. 311:560-564). Importantly, approximately 80% of urine specimens tested in clinical microbiology laboratories are considered negative (i.e. bacterial count of less than 10⁷ CFU/L; Table 3). Urine specimens found positive by culture are further characterized using standard biochemical tests to identify the bacterial pathogen and are also tested for susceptibility to antibiotics. The biochemical and susceptibility testing normally require 18-24 hours of incubation.

Accurate and rapid urine screening methods for bacterial pathogens would allow a faster identification of negative specimens and a more efficient treatment and care management of patients. Several rapid identification methods (UriscreenTM, UTIscreenTM, Flash TrackTM DNA probes and others) have been compared to slower standard biochemical methods, which are based on culture of the bacterial pathogens. Although much faster, these rapid tests showed low sensitivities and poor specificities as well as a high number of false negative and false positive results (Koening *et al.*, 1992, J. Clin. Microbiol. **30**:342-345; Pezzlo *et al.*, 1992, J. Clin. Microbiol. **30**:640-684).

Blood specimens

The blood specimens received in the microbiology laboratory are always submitted for culture. Blood culture systems may be manual, semi-automated or completely automated. The BACTECTM system (from Becton Dickinson) and the BacTAlertTM system (from Organon Teknika Corporation) are the two most widely used automated blood culture systems. These systems incubate blood culture bottles under optimal conditions for growth of most bacteria. Bacterial growth is monitored continuously to detect early positives by using highly sensitive bacterial growth detectors. Once growth is detected, a Gram stain is performed directly from the blood culture and then used to inoculate nutrient agar plates. Subsequently, bacterial identification and susceptibility testing are carried out from isolated bacterial colonies with automated systems as described previously. Blood culture bottles are normally reported as negative if no growth is detected after an incubation of 6 to 7 days. Normally, the vast majority of blood cultures are reported negative. For example, the percentage of negative blood cultures at the microbiology laboratory of the CHUL for the period February 1994-January 1995 was 93.1% (Table 3).

Other clinical samples

Upon receipt by the clinical microbiology laboratory, all body fluids other than blood and urine that are from normally sterile sites (i.e. cerebrospinal, synovial, pleural, pericardial and others) are processed for direct microscopic examination and subsequent culture. Again, most clinical samples are negative for culture (Table 3). In all these normally sterile sites, tests for the universal detection of algae, archaea, bacteria, fungi and parasites would be very useful.

Regarding clinical specimens which are not from sterile sites such as sputum or stool specimens, the laboratory diagnosis by culture is more problematic because of the contamination by the normal flora. The bacterial or fungal pathogens potentially associated with the infection are grown and separated from the colonizing microbes using selective methods and then identified as described previously. Of course, the DNA-based universal detection of bacteria would not be useful for the diagnosis of bacterial infections at these non-sterile sites. On the other hand, DNA-based assays for species or genus or family or group detection and identification as well as for the detection of antimicrobial agents resistance genes from these specimens would be very useful and would offer several advantages over classical identification and susceptibility testing methods.

DNA-based assays with any specimen

There is an obvious need for rapid and accurate diagnostic tests for the detection and identification of algae, archaea, bacteria, fungi and parasites directly from clinical specimens. DNA-based technologies are rapid and accurate and offer a great potential to improve the diagnosis of infectious diseases (Persing *et al.*, 1993, Diagnostic Molecular Microbiology: Principles and Applications, American Society for Microbiology, Washington, D.C.; Bergeron and Ouellette, 1995, Infection 23:69-72; Bergeron and Ouellette, 1998, J Clin Microbiol. 36:2169-72). The DNA probes and amplification primers which are objects of the present invention are applicable for the detection and identification of algae, archaea, bacteria, fungi, and parasites directly from any clinical specimen such as blood,

urine, sputum, cerebrospinal fluid, pus, genital and gastro-intestinal tracts, skin or any other type of specimens (Table 3). These assays are also applicable to detection from microbial cultures (e.g. blood cultures, bacterial or fungal colonies on nutrient agar, or liquid cell cutures in nutrient broth). The DNA-based tests proposed in this invention are superior in terms of both rapidity and accuracy to standard biochemical methods currently used for routine diagnosis from any clinical specimens in microbiology laboratories. Since these tests can be performed in one hour or less, they provide the clinician with new diagnostic tools which should contribute to a better management of patients with infectious diseases. Specimens from sources other than humans (e.g. other primates, birds, plants, mammals, farm animals, livestock, food products, environment such as water or soil, and others) may also be tested with these assays.

A high percentage of culture-negative specimens

Among all the clinical specimens received for routine diagnosis, approximately 80% of urine specimens and even more (around 95%) for other types of normally sterile clinical specimens are negative for the presence of bacterial pathogens (Table 3). It would also be desirable, in addition to identify bacteria at the species or genus or family or group level in a given specimen, to screen out the high proportion of negative clinical specimens with a DNA-based test detecting the presence of any bacterium (i.e. universal bacterial detection). As disclosed in the present invention, such a screening test may be based on DNA amplification by PCR of a highly conserved genetic target found in all bacteria. Specimens negative for bacteria would not be amplified by this assay. On the other hand, those that are positive for any bacterium would give a positive amplification signal. Similarly, highly conserved genes of fungi and parasites could serve not only to identify particular species or genus or family or group but also to detect the presence of any fungi or parasite in the specimen.

WO 01/23604 PCT/CA00/01150 Towards the development of rapid DNA-based diagnostic tests

A rapid diagnostic test should have a significant impact on the management of infections. DNA probe and DNA amplification technologies offer several advantages over conventional methods for the identification of pathogens and antimicrobial agents resistance genes from clinical samples (Persing et al., 1993, Diagnostic Molecular Microbiology: Principles and Applications, American Society for Microbiology, Washington, D.C.; Ehrlich and Greenberg, 1994, PCRbased Diagnostics in Infectious Disease, Blackwell Scientific Publications, Boston, MA). There is no need for culture of the pathogens, hence the organisms can be detected directly from clinical samples, thereby reducing the time associated with the isolation and identification of pathogens. Furthermore, DNA-based assays are more accurate for microbial identification than currently used phenotypic identification systems which are based on biochemical tests and/or microscopic examination. Commercially available DNA-based technologies are currently used in clinical microbiology laboratories, mainly for the detection and identification of fastidious bacterial pathogens such as Mycobacterium tuberculosis, Chlamydia trachomatis, Neisseria gonorrhoeae as well as for the detection of a variety of viruses (Tang Y. and Persing D. H., Molecular detection and identification of microorganisms, In: P. Murray et al., 1999, Manual of Clinical Microbiology, ASM press, 7th edition, Washington D.C.). There are also other commercially available DNA-based assays which are used for culture confirmation assays.

Others have developed DNA-based tests for the detection and identification of bacterial pathogens which are objects of the present invention, for example: Staphylococcus sp. (US patent serial no. 5,437,978), Neisseria sp. (US patent serial no. 5,162,199 and European patent serial no. 0,337,896,131) and Listeria monocytogenes (US patent serial nos. 5,389,513 and 5,089,386). However, the diagnostic tests described in these patents are based either on rRNA genes or on genetic targets different from those described in the present invention. To our knowledge there are only four patents published by others mentioning the use of

any of the four highly conserved gene targets described in the present invention for diagnostic purposes (PCT international publication number WO92/03455 and WO00/14274, European patent publication number 0 133 671 B1, and European patent publication number 0 133 288 A2). WO92/03455 is focused on the inhibition of Candida species for therapeutic purposes. It describes antisense oligonucleotide probes hybridizing to Candida messenger RNA. Two of the numerous mRNA proposed as targets are coding for translation elongation factor 1 (tef1) and the beta subunit of ATPase. DNA amplification or hybrization are not under the scope of their invention and although diagnostic use is briefly mentioned in the body of the application, no specific claim is made regarding diagnostics. WO00/14274 describes the use of bacterial recA gene for identification and speciation of bacteria of the Burkholderia cepacia complex. Specific claims are made on a method for obtaining nucleotide sequence information for the recA gene from the target bacteria and a following comparison with a standard library of nucleotide sequence information (claim 1), and on the use of PCR for amplification of the recA gene in a sample of interest (claims 4 to 7, and 13). However, the use of a discriminatory restriction enzyme in a RFLP procedure is essential to fulfill the speciation and WO00/14274 did not mention that multiple recA probes could be used simultaneously. Patent EP 0 133 288 A2 describes and claims the use of bacterial tuf (and fus) sequence for diagnostics based on hybridization of a tuf (or fus) probe with bacterial DNA. DNA amplification is not under the scope of EP 0 133 288 A2. Nowhere it is mentioned that multiple tuf (or fus) probes could be used simultaneously. No mention is made regarding speciation using tuf (or fus) DNA nucleic acids and/or sequences. The sensitivities of the tuf hybrizations reported are $1x10^6$ bacteria or 1-100 ng of DNA. This is much less sensitive than what is achieved by our assays using nucleic acid amplification technologies.

Although there are phenotypic identification methods which have been used for more than 125 years in clinical microbiology laboratories, these methods do not provide information fast enough to be useful in the initial management of patients.

There is a need to increase the speed of the diagnosis of commonly encountered bacterial, fungal and parasitical infections. Besides being much faster, DNA-based diagnostic tests are more accurate than standard biochemical tests presently used for diagnosis because the microbial genotype (e.g. DNA level) is more stable than the phenotype (e.g. physiologic level).

Bacteria, fungi and parasites encompass numerous well-known microbial pathogens. Other microorganisms could also be pathogens or associated with human diseases. For example, achlorophylious algae of the *Prototheca* genus can infect humans. Archae, especially methanogens, are present in the gut flora of humans (Reeve, J.H., 1999, J. Bacteriol. 181:3613-3617). However, methanogens have been associated to pathologic manifestations in the colon, vagina, and mouth (Belay *et al.*, 1988, Appl. Enviro. Microbiol. 54:600-603; Belay *et al.*, 1990, J. Clin. Microbiol. 28:1666-1668; Weaver *et al.*, 1986, Gut 27:698-704).

In addition to the identification of the infectious agent, it is often desirable to identify harmful toxins and/or to monitor the sensitivity of the microorganism to antimicrobial agents. As revealed in this invention, genetic identification of the microorganism could be performed simultaneously with toxin and antimicrobial agents resistance genes.

Knowledge of the genomic sequences of algal, archaeal, bacterial, fungal and parasitical species continuously increases as testified by the number of sequences available from public databases such as GenBank. From the sequences readily available from those public databases, there is no indication therefrom as to their potential for diagnostic purposes. For determining good candidates for diagnostic purposes, one could select sequences for DNA-based assays for (i) the species-specific detection and identification of commonly encountered bacterial, fungal and parasitical pathogens, (ii) the genus-specific detection and identification of commonly encountered bacterial, fungal or parasitical pathogens, (iii) the family-specific detection and identification of commonly encountered bacterial, fungal or parasitical pathogens, (iv) the group-specific detection and identification of commonly encountered bacterial, fungal or parasitical pathogens, (v) the

universal detection of algal, archaeal, bacterial, fungal or parasitical pathogens, and/or (vi) the specific detection and identification of antimicrobial agents resistance genes, and/or (vii) the specific detection and identification of bacterial toxin genes. All of the above types of DNA-based assays may be performed directly from any type of clinical specimens or from a microbial culture.

In our assigned U.S. patent 6,001,564 and our WO98/20157 patent publication, we described DNA sequences suitable for (i) the species-specific detection and identification of clinically important bacterial pathogens, (ii) the universal detection of bacteria, and (iii) the detection of antimicrobial agents resistance genes.

The WO98/20157 patent publication describes proprietary tuf DNA sequences as well as tuf sequences selected from public databases (in both cases, fragments of at least 100 base pairs), as well as oligonucleotide probes and amplification primers derived from these sequences. All the nucleic acid sequences described in that patent publication can enter in the composition of diagnostic kits or products and methods capable of a) detecting the presence of bacteria and fungi b) detecting specifically at the species, genus, family or group levels, the presence of bacteria and fungi and antimicrobial agents resistance genes associated with these pathogens. However, these methods and kits need to be improved, since the ideal kit and method should be capable of diagnosing close to 100% of microbial pathogens and associated antimicrobial agents resistance genes and toxins genes. For example, infections caused by Enterococcus faecium have become a clinical problem because of its resistance to many antibiotics. Both the detection of these bacteria and the evaluation of their resistance profiles are desirable. Besides that, novel DNA sequences (probes and primers) capable of recognizing the same and other microbial pathogens or the same and additional antimicrobial agents resistance genes are also desirable to aim at detecting more target genes and complement our earlier patent applications.

The present invention improves the assigned application by disclosing new proprietary tuf nucleic acids and/or sequences as well as describing new ways to

obtain *tuf* nucleic acids and/or sequences. In addition we disclose new proprietary *atpD* and *recA* nucleic acids and/or sequences. In addition, new uses of *tuf*, *atpD* and *recA* DNA nucleic acids and/or sequences selected from public databases (Table 11) are disclosed.

Highly conserved genes for identification and diagnostics

Highly conserved genes are useful for identification of microorganisms. For bacteria, the most studied genes for identification of microorganisms are the universally conserved ribosomal RNA genes (rRNA). Among those, the principal targets used for identification purposes are the small subunit (SSU) ribosomal 16S rRNA genes (in prokaryotes) and 18S rRNA genes (in eukaryotes) (Relman and Persing, Genotyping Methods for Microbial Identification, *In*: D.H. Persing, 1996, PCR Protocols for Emerging Infectious Diseases, ASM Press, Washington D.C.). The rRNA genes are also the most commonly used targets for universal detection of bacteria (Chen *et al.*, 1988, FEMS Microbiol. Lett. **57**:19-24; McCabe *et al.*, 1999, Mol. Genet. Metabol. **66**:205-211) and fungi (Van Burik *et al.*, 1998, J. Clin. Microbiol. **36**:1169-1175).

However, it may be difficult to discriminate between closely related species when using primers derived from the 16S rRNA. In some instances, 16S rRNA sequence identity may not be sufficient to guarantee species identity (Fox et al., 1992, Int. J. Syst. Bacteriol. 42:166-170) and it has been shown that inter-operon sequence variation as well as strain to strain variation could undermine the application of 16S rRNA for identification purposes (Clayton et al., 1995, Int. J. Syst. Bacteriol. 45:595-599). The heat shock proteins (HSP) are another family of very conserved proteins. These ubiquitous proteins in bacteria and eukaryotes are expressed in answer to external stress agents. One of the most described of these HSP is HSP 60. This protein is very conserved at the amino acid level, hence it has been useful for phylogenetic studies. Similar to 16S rRNA, it would be difficult to

discriminate between species using the HSP 60 nucleotide sequences as a diagnostic tool. However, Goh et al. identified a highly conserved region flanking a variable region in HSP 60, which led to the design of universal primers amplifying this variable region (Goh et al., US patent serial no. 5,708,160). The sequence variations in the resulting amplicons were found useful for the design of species-specific assays.

SUMMARY OF THE INVENTION

It is an object of the present invention to provide a specific, ubiquitous and sensitive method using probes and/or amplification primers for determining the presence and/or amount of nucleic acids:

- from any algal, archaeal, bacterial, fungal or parasitical species in any sample suspected of containing said nucleic acids, and optionally,
- from specific microbial species or genera selected from the group consisting of the species or genera listed in Table 4, and optionally,
- from an antimicrobial agents resistance gene selected from the group consisting of the genes listed in Table 5, and optionally,
- from a toxin gene selected from the group consisting of the genes listed in Table 6,

wherein each of said nucleic acids or a variant or part thereof comprises a selected target region hybridizable with said probes or primers;

said method comprising the steps of contacting said sample with said probes or primers and detecting the presence and/or amount of hybridized probes or amplified products as an indication of the presence and/or amount of said any WO 01/23604 PCT/CA00/01150 microbial species, specific microbial species or genus or family or group and antimicrobial agents resistance gene and/or toxin gene.

In a specific embodiment, a similar method directed to each specific microbial species or genus or family or group detection and identification, antimicrobial agents resistance genes detection, toxin genes detection, and universal bacterial detection, separately, is provided.

In a more specific embodiment, the method makes use of DNA fragments from conserved genes (proprietary sequences and sequences obtained from public databases), selected for their capacity to sensitively, specifically and ubiquitously detect the targeted algal, archaeal, bacterial, fungal or parasitical nucleic acids.

In a particularly preferred embodiment, oligonucleotides of at least 12 nucleotides in length have been derived from the longer DNA fragments, and are used in the present method as probes or amplification primers. To be a good diagnostic candidate, an oligonucleotide of at least 12 nucleotides should be capable of hybridizing with nucleic acids from given microorganism(s), and with substantially all strains and representatives of said microorganism(s); said oligonucleotide being species-, or genus-, or family-, or group-specific or universal.

In another particularly preferred embodiment, oligonucleotides primers and probes of at least 12 nucleotides in length are designed for their specificity and ubiquity based upon analysis of our databases of *tuf*, *atpD* and *recA* sequences. These databases are generated using both proprietary and public sequence information. Altogether, these databases form a sequence repertory useful for the design of primers and probes for the detection and identification of algal, archaeal, bacterial, fungal and parasitical microorganisms. The repertory can also be subdivided into subrepertories for sequence analysis leading to the design of various primers and probes.

The *tuf*, *atpD* and *recA* sequences databases as a product to assist the design of oligonucleotides primers and probes for the detection and identification of algal, archaeal, bacterial, fungal and parasitical microorganisms are also covered.

The proprietary oligonucleotides (probes and primers) are also another object of this invention.

Diagnostic kits comprising probes or amplification primers such as those for the detection of a microbial species or genus or family or phylum or group selected from the following list consisting of Abiotrophia adiacens, Acinetobacter baumanii, Actinomycetae, Bacteroides, Cytophaga and Flexibacter phylum, Bacteroides fragilis, Bordetella pertussis, Bordetella sp., Campylobacter jejuni and C. coli, Candida albicans, Candida dubliniensis, Candida glabrata, Candida guilliermondii, Candida krusei, Candida lusitaniae, Candida parapsilosis, Candida tropicalis, Candida zeylanoides, Candida sp., Chlamydia pneumoniae, Chlamydia trachomatis, Clostridium sp., Corynebacterium sp., Crypococcus neoformans, Cryptococcus sp., Cryptosporidium parvum, Entamoeba sp., Enterobacteriaceae group, Enterococcus casseliflavus-flavescens-gallinarum group, Enterococcus faecalis, Enterococcus faecium, Enterococcus gallinarum, Enterococcus sp., Escherichia coli and Shigella sp. group, Gemella sp., Giardia sp., Haemophilus influenzae, Klebsiella pneumoniae, Legionella pneumophila, Legionella sp., Leishmania sp., Mycobacteriaceae family, Mycoplasma pneumoniae, Neisseria gonorrhoeae, platelets contaminants group (see Table 14), Pseudomonas aeruginosa, Pseudomonads group, Staphylococcus aureus, Staphylococcus epidermidis, Staphylococcus haemolyticus, Staphylococcus hominis, Staphylococcus saprophyticus, Staphylococcus sp., Streptococcus agalactiae, Streptococcus pneumoniae, Streptococcus pyogenes, Streptococcus sp., Trypanosoma brucei, Trypanosoma cruzi, Trypanosoma sp., Trypanosomatidae family, are also objects of the present invention.

Diagnostic kits further comprising probes or amplification primers for the detection of an antimicrobial agents resistance gene selected from the group listed in Table 5 are also objects of this invention.

Diagnostic kits further comprising probes or amplification primers for the detection of a toxin gene selected from the group listed in Table 6 are also objects of this invention.

Diagnostic kits further comprising probes or amplification primers for the detection of any other algal, archaeal, bacterial, fungal or parasitical species than those specifically listed herein, comprising or not comprising those for the detection of the specific microbial species or genus or family or group listed above, and further comprising or not comprising probes and primers for the antimicrobial agents resistance genes listed in Table 5, and further comprising or not comprising probes and primers for the toxin genes listed in Table 6 are also objects of this invention.

In a preferred embodiment, such a kit allows for the separate or the simultaneous detection and identification of the above-listed microbial species or genus or family or group; or universal detection of algae, archaea, bacteria, fungi or parasites; or antimicrobial agents resistance genes; or toxin genes; or for the detection of any microorganism (algae, archaea, bacteria, fungi or parasites).

In the above methods and kits, probes and primers are not limited to nucleic acids and may include, but are not restricted to analogs of nucleotides such as: inosine, 3-nitropyrrole nucleosides (Nichols *et al.*, 1994, Nature **369**:492-493), Linked Nucleic Acids (LNA) (Koskin *et al.*, 1998, Tetrahedron **54**:3607-3630), and Peptide Nucleic Acids (PNA) (Egholm *et al.*, 1993, Nature **365**:566-568).

In the above methods and kits, amplification reactions may include but are not restricted to: a) polymerase chain reaction (PCR), b) ligase chain reaction (LCR), c) nucleic acid sequence-based amplification (NASBA), d) self-sustained sequence replication (3SR), e) strand displacement amplification (SDA), f) branched DNA signal amplification (bDNA), g) transcription-mediated amplification (TMA), h) cycling probe technology (CPT), i) nested PCR, j) multiplex PCR, k) solid phase amplification (SPA), l) nuclease dependent signal amplification (NDSA), m) rolling circle amplification technology (RCA), n) Anchored strand displacement amplification, o) Solid-phase (immobilized) rolling circle amplification.

In the above methods and kits, detection of the nucleic acids of target genes may include real-time or post-amplification technologies. These detection

transfer (FRET)-based methods such as adjacent hybridization to FRET probes (including probe-probe and probe-primer methods), TaqMan, Molecular Beacons, scorpions, nanoparticle probes and Sunrise (Amplifluor). Other detection methods include target genes nucleic acids detection via immunological methods, solid phase hybridization methods on filters, chips or any other solid support, whether the hybridization is monitored by fluorescence, chemiluminescence, potentiometry, mass spectrometry, plasmon resonance, polarimetry, colorimetry, or scanometry. Sequencing, including sequencing by dideoxy termination or sequencing by hybridization, e.g. sequencing using a DNA chip, is another possible method to detect and identify the nucleic acids of target genes.

In a preferred embodiment, a PCR protocol is used for nucleic acid amplification, in diagnostic method as well as in method of construction of a repertory of nucleic acids and deduced sequences.

In a particularly preferred embodiment, a PCR protocol is provided, comprising, an initial denaturation step of 1-3 minutes at 95 °C, followed by an amplification cycle including a denaturation step of one second at 95 °C and an annealing step of 30 seconds at 45-65°C, without any time allowed specifically for the elongation step. This PCR protocol has been standardized to be suitable for PCR reactions with most selected primer pairs, which greatly facilitates the testing because each clinical sample can be tested with universal, species-specific, genus-specific, antimicrobial agents resistance gene and toxin gene PCR primers under uniform cycling conditions. Furthermore, various combinations of primer pairs may be used in multiplex PCR assays.

It is also an object of the present invention that *tuf*, *atpD* and *recA* sequences could serve as drug targets and these sequences and means to obtain them revealed in the present invention can assist the screening, design and modeling of these drugs.

It is also an object of the present invention that tuf, atpD and recA sequences could serve for vaccine purposes and these sequences and means to obtain them

revealed in the present invention can assist the screening, design and modeling of these vaccines.

We aim at developing a universal DNA-based test or kit to screen out rapidly samples which are free of algal, archaeal, bacterial, fungal or parasitical cells. This test could be used alone or combined with more specific identification tests to detect and identify the above algal and/or archaeal and/or bacterial and/or fungal and/or parasitical species and/or genera and/or family and/or group and to determine rapidly the bacterial resistance to antibiotics and/or presence of bacterial toxins. Although the sequences from the selected antimicrobial agents resistance genes are available from public databases and have been used to develop DNAbased tests for their detection, our approach is unique because it represents a major improvement over current diagnostic methods based on bacterial cultures. Using an amplification method for the simultaneous or independent or sequential microbial detection-identification and antimicrobial resistance genes detection, there is no need for culturing the clinical sample prior to testing. Moreover, a modified PCR protocol has been developed to detect all target DNA sequences in approximately one hour under uniform amplification conditions. This procedure should save lives by optimizing treatment, should diminish antimicrobial agents resistance because less antibiotics will be prescribed, should reduce the use of broad spectrum antibiotics which are expensive, decrease overall health care costs by preventing or shortening hospitalizations, and side effects of drugs, and decrease the time and costs associated with clinical laboratory testing.

In another embodiment, sequence repertories and ways to obtain them for other gene targets are also an object of this invention, such is the case for the *hexA* nucleic acids and/or sequences of Streptococci.

In yet another embodiment, for the detection of mutations associated with antibiotic resistance genes, we built repertories to distinguish between point mutations reflecting only gene diversity and point mutations involved in resistance. Such repertories and ways to obtain them for pbp1a, pbp2b and pbp2x genes of sensitive and penicillin-resistant Streptoccoccus pneumoniae and also for gyrA and

parC gene fragments from various bacterial species are also an object of the present invention.

The diagnostic kits, primers and probes mentioned above can be used to identify algae, archaea, bacteria, fungi, parasites, antimicrobial agents resistance genes and toxin genes on any type of sample, whether said diagnostic kits, primers and probes are used for *in vitro* or *in situ* applications. The said samples may include but are not limited to: any clinical sample, any environment sample, any microbial culture, any microbial colony, any tissue, and any cell line.

It is also an object of the present invention that said diagnostic kits, primers and probes can be used alone or in conjunction with any other assay suitable to identify microorganisms, including but not limited to: any immunoassay, any enzymatic assay, any biochemical assay, any lysotypic assay, any serological assay, any differential culture medium, any enrichment culture medium, any selective culture medium, any specific assay medium, any identification culture medium, any enumeration cuture medium, any cellular stain, any culture on specific cell lines, and any infectivity assay on animals.

In the methods and kits described herein below, the oligonucleotide probes and amplification primers have been derived from larger sequences (i.e. DNA fragments of at least 100 base pairs). All DNA fragments have been obtained either from proprietary fragments or from public databases. DNA fragments selected from public databases are newly used in a method of detection according to the present invention, since they have been selected for their diagnostic potential.

In another embodiment, the amino acid sequences translated from the repertory of *tuf*, *atpD* and *recA* nucleic acids and/or sequences are also an object of the present invention.

It is clear to the individual skilled in the art that other oligonucleotide sequences appropriate for (i) the universal detection of algae, archaea, bacteria, fungi or parasites, (ii) the detection and identification of the above microbial species or genus or family or group, and (iii) the detection of antimicrobial agents resistance genes, and (iv) the detection of toxin genes, other than those listed in

Annexes I to III, XXI to XXII, XXXII to XXXVII, XXXIX to XLI, and XLIII to LIV may also be derived from the proprietary fragments or selected public database sequences. For example, the oligonucleotide primers or probes may be shorter or longer than the ones chosen; they may also be selected anywhere else in the proprietary DNA fragments or in the sequences selected from public databases; they may be also variants of the same oligonucleotide. If the target DNA or a variant thereof hybridizes to a given oligonucleotide, or if the target DNA or a variant thereof can be amplified by a given oligonucleotide PCR primer pair, the converse is also true; a given target DNA may hybridize to a variant oligonucleotide probe or be amplified by a variant oligonucleotide PCR primer. Alternatively, the oligonucleotides may be designed from any DNA fragment sequences for use in amplification methods other than PCR. Consequently, the core of this invention is the identification of universal, species-specific, genus-specific, family-specific, group-specific, resistance gene-specific, toxin gene-specific genomic or non-genomic DNA fragments which are used as a source of specific and ubiquitous oligonucleotide probes and/or amplification primers. Although the selection and evaluation of oligonucleotides suitable for diagnostic purposes requires much effort, it is quite possible for the individual skilled in the art to derive, from the selected DNA fragments, oligonucleotides other than the ones listed in Annexes I to III, XXI to XXII, XXXII to XXXVII, XXXIX to XLI, and XLIII to LIV which are suitable for diagnostic purposes. When a proprietary fragment or a public databases sequence is selected for its specificity and ubiquity, it increases the probability that subsets thereof will also be specific and ubiquitous.

Since a high percentage of clinical specimens are negative for bacteria (Table 3), DNA fragments having a high potential for the selection of universal oligonucleotide probes or primers were selected from proprietary and public database sequences. The amplification primers were selected from genes highly conserved in algae, archaea, bacteria, fungi and parasites, and are used to detect the presence of any algal, archaeal, bacterial, fungal or parasitical pathogen in clinical specimens in order to determine rapidly whether it is positive or negative for algae,

archaea, bacteria, fungi or parasites. The selected genes, designated *tuf*, *fus*, *atpD* and *recA*, encode respectively 2 proteins (elongation factors Tu and G) involved in the translational process during protein synthesis, a protein (beta subunit) responsible for the catalytic activity of proton pump ATPase and a protein responsible for the homologous recombination of genetic material. The alignments of *tuf*, *atpD* and *recA* sequences used to derive the universal primers include both proprietary and public database sequences. The universal primer strategy allows the rapid screening of the numerous negative clinical specimens (around 80% of the specimens received, see Table 3) submitted for microbiological testing.

Table 4 provides a list of the archaeal, bacterial, fungal and parasitical species for which *tuf* and/or *atpD* and/or *recA* nucleic acids and/or sequences are revealed in the present invention. Tables 5 and 6 provide a list of antimicrobial agents resistance genes and toxin genes selected for diagnostic purposes. Table 7 provides the origin of *tuf*, *atpD* and *recA* nucleic acids and/or sequences listed in the sequence listing. Tables 8-10 and 12-14 provide lists of species used to test the specificity, ubiquity and sensitivity of some assays described in the examples. Table 11 provides a list of microbial species for which *tuf* and/or *atpD* and/or *recA* sequences are available in public databases. Table 15 lists the microorganisms identified by commercial systems. Tables 16-18 are part of Example 42, whereas Tables 19-20 are part of Example 43. Tables 21-22 illustrate Example 44, whereas Tables 23-25 illustrate Example 45.

In accordance with the present invention is provided a method for generating a repertory of nucleic acids of *tuf*, *fus*, *atpD* and/or *recA* genes from which are derived probes or primers, or both, useful for the detection of one, more than one related microorganisms, or substantially all microorganisms of a group selected from algae, archaea, bacteria, fungi and parasites, which comprises the step of:

amplifying the nucleic acids of a plurality of determined algal, archaeal, bacterial, fungal and parasitical species with any combination of the primer pairs defined in SEQ ID NOs.: 558-561, 562-574, 636-655, 664, 681-683, 696-697, 699-700, 708, 812-815, 911-917, 919-922, 935-938, 1203-1207, 1212-1213, 1221-1229, 1605-1606, 1974-1984, 1999- 2003, 2282-2285.

The terms "related microorganisms" are intended to cover microorganisms that share a common evolutive profile up to the speciation e.g. those that belong to a species, a genus, a family or a phyllum. The same terms are also intended to cover a group of different species that are grouped for a specific reason, for example, because they all have a common host tissue or cell. In one specific example, a group of microorganims potentially found in platelet preparations are grouped together and are considered "related" organisms for the purpose of their simultaneous detection in that particular type of sample.

The repertories *per se* of nucleic acids and of sequences derived therefrom are also provided, as well as "gene banks" comprising these repertories.

For generating sequences of probes or primers, the above method is reproduced or one may start from the sequence repertory or gene bank itself, and the following steps are added:

- aligning a subset of nucleic acid sequences of said repertory,
- locating nucleic acid stretches that are present in the nucleic acids of strains or representatives of said one, more than one related microorganisms, or substantially all microorganisms of said group, and not present in the nucleic acid sequences of other microorganisms, and

deriving consensus nucleic acid sequ nces useful as probes or primers from said stretches.

Once the sequences of probes or primers are designed, they are converted into real molecules by nucleic acid synthesis.

From the above methods and resulting repertories, probes and primers for the universal detection of any one of alga, archaeon, bacterium, fungus and parasite are obtainable.

More specifically, the following probes or primers having the sequence defined in SEQ ID NOs.: 543, 556-574, 636-655, 658-661, 664, 681-683, 694, 696, 697, 699, 700, 708, 812-815, 911-917, 919-922, 935-938, 1203-1207, 1212-1213, 1221-1229, 1605-1606, 1974-1984, 1999-2000, 2282-2285 or any variant of at least 12 nucleotides capable of hybridizing with the targeted microorganism(s) and these sequences and a diagnostic method using the same are provided.

Further, probes or primers having specific and ubiquitous properties for the detection and identification of any one of an algal, archaeal, bacterial, fungal and parasitital species, genus, family and group are also designed and derived from the same methods and repertories.

More specifically, are provided definite probes or primers having specific and ubiquitous properties for the detection and identification of microorganisms.

Indeed, a general method is provided for detecting the presence in a test sample of any microorganism that is an alga, archaeum, bacterium, fungus or parasite, which comprises:

a) putting in contact any test sample *tuf* or *atpD* or *recA* sequences and nucleic acid primers and/or probes, said primers and/or probes having be n selected to be sufficiently complementary to hybridize to

one or more *tuf* or *atpD* or *recA* sequences that are specific to said microorganism:

- b) allowing the primers and/or probes and any test sample *tuf* or *atpD* or *recA* sequences to hybridize under specified conditions such as said primers and/or probes hybridize to the *tuf* or *atpD* or *recA* sequences of said microorganism and does not delectably hybridize to *tuf* or *atpD* or *recA* sequences from other microorganisms; and,
- c) testing for hybridization of said primers and/or probes to any test sample *tuf* or *atpD* or *recA* sequences.

In the latter, step c) is based on a nucleic acid target amplification method, or on a signal amplification method.

The terms "sufficiently complementary" cover perfect and imperfect complementarity.

In addition to the universal or the specific detection and/or identification of microorganisms, the simultaneous detection of antimicrobial agent resistance gene or of a toxin gene is provided in compositions of matter as well as in diagnostic methods. Such detection is brought by using probes or primers having at least 12 nucleotides in length capable of hybridizing with an antimicrobial agent resistance gene and/or toxin gene, a definite set thereof being particularly provided.

Of course, any propriatory nucleic acid and nucleotide sequence derived therefrom, and any variant of at least 12 nucleotides capable of a selective hybridization with the following nucleic acids are within the scope of this invention as well as derived recombinant vectors and hosts:

SEQ ID NOs.: 1-73, 75-241, 399-457, 498-529, 612-618, 621-624, 675, 677, 717-736, 779-792, 840-855, 865, 868-888, 897-910, 932, 967-989 992, 1266-1297, 1518-1526, 1561-1575, 1578-1580, 1662-1664, 1666-1667, 1669-1670, 1673-1683, 1685-1689, 1786-1843, 1874-1881, 1956-1960, 2183-2185, 2187-2188, 2193-2201, 2214-2249, 2255-2272, which are all *tuf* sequences;

SEO ID NOs.: 242-270, 272-398, 458-497, 530-538, 663, 667, 673-676, 678-680, 737-778, 827-832, 834-839, 856-862, 866-867, 889-896, 929-931, 941-966, 1245-1254, 1256-1265, 1527, 1576-1577, 1600-1604,1638-1647, 1649-1660, 1671, 1684, 1844-1848, 1849-1865, 2189-2192, which are all *atpD* sequences;

SEQ ID NOs.: 990-991, 1003, 1288-1289, 1714, 1756-1763, 1866-1873 and 2202-2212, which are all *recA* sequences; and

SEQ ID NOs.: 1004-1075, 1255, 1607-1608, 1648, 1764-1785, 2013-2014, 2056-2064, 2273-2280, which are antimicrobial agent resistance or toxin gene sequences found to be suitable for the detection and identification of microbial species.

To complement the following repertories, another one comprising *hexA* nucleic acids and derived sequences have been construed through amplification of nucleic acids of any streptococcal species with any combination of primers SEO ID NOs.: 1179, 1181, 1182 and 1184 to 1191. From this particular repertory, primers and/or probes for detecting *Streptococcus pneumoniae* have been designed and obtained. Particularly, a nucleic acid sequence of at least 12 nucleotides capable of hybridizing with *Streptococcus pneumoniae* and with any one of SEQ ID NOs.: 1184 to 1187 or with SEQ ID NOs.: 1179, 1180, 1181 or 1182 are provided.

The remarkable sequence diversity of nucleic acids that encode proteins also provides diversity of peptide sequences which constitute another repertory that is also within the scope of this invention. From the protein and nucleic acid sequence repertories is derived a use therefrom for the design of a therapeutic agent effective against a target microorganism, for example, an antibiotic, a vaccine or a genic therapeutic agent.

Due to the constant evolution in the diagnostic methods, here is finally provided a method for the identification of a microorganism in a test sample, comprising the steps of:

a) obtaining a nucleic acid sequence from a *tuf*, *fus*, *atpD*, and/or *recA* genes of said microorganisms, and

b) comparing said nucleic acid sequence with the nucleic acid sequences of a bank as defined in claim 5, said repertory comprising a nucleic acid sequence obtained from the nucleic acids of said microorganism, whereby said microorganism is identify when there is a match between the sequences.

In this method, any way by which the specified given sequence is obtained is contemplated, and this sequence is simply compared to the sequences of a bank or a repertory. If the comparison results in a match, e.g. if bank comprises the nucleic acid sequence of interest, the identification of the microorganism is provided.

DETAILED DESCRIPTION OF THE INVENTION

HIGHLY CONSERVED GENES AND THEIR USE TO GENERATE SPECIESSPECIFIC, GENUS-SPECIFIC, FAMILY-SPECIFIC, GROUP-SPECIFIC AND
UNIVERSAL NUCLEIC ACID PROBES AND AMPLIFICATION PRIMERS TO
RAPIDLY DETECT AND IDENTIFY ALGAL, ARCHAEAL, BACTERIAL,
FUNGAL AND PARASITICAL MICROORGANISMS FROM CLINICAL
SPECIMENS FOR DIAGNOSIS

The present inventors reasoned that comparing the published *Haemophilus* influenzae and *Mycoplasma genitalium* genomes and searching for conserved genes could provide targets to develop useful diagnostic primers and probes. This sequence comparison is highly informative as these two bacteria are distantly related and most genes present in the minimal genome of *M. genitalium* are likely to be present in every bacterium. Therefore genes conserved between these two bacteria are likely to be conserved in all other bacteria.

Following the genomic comparison, it was found that several protein-coding genes were conserved in evolution. Highly conserved proteins included the translation elongation factors G (EF-G) and Tu (EF-Tu) and the β subunit of F0F1 type ATP-synthase, and to a lesser extent, the RecA recombinase. These four proteins coding genes were selected amongst the 20 most conserved genes on the basis that they all possess at least two highly conserved regions suitable for the design of universal amplification and sequencing primers. Moreover, within the fragment amplified by these primers, highly conserved and more variable regions are also present hence suggesting it might be possible to rapidly obtain sequence information from various microbial species to design universal as well as species, genus-, family-, or group-specific primers and probes of potential use for the detection and identification and/or quantification of microorganisms.

Translation elongation factors are members of a family of GTP-binding proteins which intervene in the interactions of tRNA molecules with the ribosome machinery during essential steps of protein synthesis. The role of elongation factor Tu is to facilitate the binding of aminoacylated tRNA molecules to the A site of the ribosome. The eukaryotic, archaeal (archaebacterial) and algal homolog of EF-Tu is called elongation factor 1 alpha (EF-1α). All protein synthesis factors originated from a common ancestor via gene duplications and fusions (Cousineau *et al.*, 1997, J. Mol. Evol. **45**:661-670). In particular, elongation factor G (EF-G), although having a functional role in promoting the translocation of aminoacyl-tRNA molecules from the A site to the P site of the ribosome, shares sequence homologies with EF-Tu and is thought to have arisen from the duplication and fusion of an ancestor of the EF-Tu gene.

In addition, EF-Tu is known to be the target for antibiotics belonging to the elfamycin's group as well as to other structural classes (Anborgh and Parmeggiani, 1991, EMBO J. 10:779-784; Luiten et al., 1992, European patent application serial No. EP 0 466 251 A1). EF-G for its part, is the target of the antibiotic fusidic acid. In addition to its crucial activities in translation, EF-Tu has chaperone-like functions in protein folding, protection against heat denaturation of proteins and interactions with unfolded proteins (Caldas et al., 1998, J. Biol. Chem 273:11478-11482). Interestingly, a form of the EF-Tu protein has been identified as a dominant component of the periplasm of Neisseria gonorrhoeae (Porcella et al., 1996, Microbiology 142:2481-2489), hence suggesting that at least in some bacterial species, EF-Tu might be an antigen with vaccine potential.

F0F1 type ATP-synthase belongs to a superfamily of proton-translocating ATPases divided in three major families: P, V and F (Nelson and Taiz, 1989, TIBS 14:113-116). P-ATPases (or E₁-E₂ type) operate via a phosphorylated intermediate and are not evolutionarily related to the other two families. V-ATPases (or V0V1 type) are present on the vacuolar and other endomembranes of eukaryotes, on the plasma membrane of archaea (archaebacteria) and algae, and also on the plasma membrane of some eubacteria especially species belonging to the order

Spirochaetales as well as to the Chlamydiaceae and Deinococcaceae families. F-ATPases (or F0F1 type) are found on the plasma membrane of most eubacteria, on the inner membrane of mitochondria and on the thylakoid membrane of chloroplasts. They function mainly in ATP synthesis. They are large multimeric enzymes sharing numerous structural and functional features with the V-ATPases. F and V-type ATPases have diverged from a common ancestor in an event preceding the appearance of eukaryotes. The β subunit of the F-ATPases is the catalytic subunit and it possesses low but significant sequence homologies with the catalytic A subunit of V-ATPases.

The translation elongation factors EF-Tu, EF-G and EF-1α, and the catalytic subunit of F or V-types ATP-synthase, are highly conserved proteins sometimes used for phylogenetic analysis and their genes are also known to be highly conserved (Iwabe *et al.*, 1989, Proc. Natl. Acad. Sci. USA 86:9355-9359, Gogarten *et al.*, 1989, Proc. Natl. Acad. Sci. USA 86:6661-6665, Ludwig *et al.*, 1993, Antonie van Leeuwenhoek 64:285-305). A recent BLAST (Altschul *et al.*, 1997, J. Mol. Biol. 215:403-410) search performed by the present inventors on the GenBank, European Molecular Biology Laboratory (EMBL), DNA Database of Japan (DDBJ) and specific genome project databases indicated that throughout bacteria, the EF-Tu and the β subunit of F0F1 type ATP-synthase genes may be more conserved than other genes that are well conserved between *H. influenzae* and *M. genitalium*.

The RecA recombinase is a multifunctional protein encoded by the *recA* gene. It plays a central role in homologous recombination, it is critical for the repair of DNA damage and it is involved in the regulation of the SOS system by promoting the proteolytic digestion of the LexA repressor. It is highly conserved in bacteria and could serve as a useful genetic marker to reconstruct bacterial phylogeny (Miller and Kokjohn, 1990, Annu. Rev. Microbiol. 44:365-394). Although RecA possesses some highly conserved sequence segments that we used to design universal primers aimed at sequencing the *recA* fragments, it is clearly not as well conserved EF-G, EF-Tu and β subunit of F0F1 type ATP-synthase.

Hence, RecA may not be optimal for universal detection of bacteria with high sensitivity but it was chosen because preliminary data indicated that EF-G, EF-Tu and β subunit of F0F1 type ATP-synthase may sometimes be too closely related to find specific primer pairs that could discriminate between certain very closely related species and genera. While RecA, EF-G, EF-Tu and β subunit of F0F1 type ATP-synthase genes, possesses highly conserved regions suitable for the design of universal sequencing primers, the less conserved region between primers should be divergent enough to allow species-specific and genus-specific primers in those cases.

Thus, as targets to design primers and probes for the genetic detection of microorganisms, the present inventors have focused on the genes encoding these four proteins: tuf, the gene for elongation factor Tu (EF-Tu); fus, the gene for the elongation factor G (EF-G); atpD, the gene for β subunit of F0F1 type ATPsynthase; and recA, the gene encoding the RecA recombinase. In several bacterial genomes tuf is often found in two highly similar duplicated copies named tufA and tufB (Filer and Furano, 1981, J. Bacteriol. 148:1006-1011, Sela et al., 1989, J. Bacteriol. 171:581-584). In some particular cases, more divergent copies of the tuf genes can exist in some bacterial species such as some actinomycetes (Luiten et al. European patent application publication No. EP 0 446 251 A1; Vijgenboom et al., 1994, Microbiology 140:983-998) and, as revealed as part of this invention, in several enterococcal species. In several bacterial species, tuf is organized in an operon with its homolog gene for the elongation factor G (EF-G) encoded by the fusA gene (Figure 3). This operon is often named the str operon. The tuf, fus, atpD and recA genes were chosen as they are well conserved in evolution and have highly conserved stretches as well as more variable segments. Moreover, these four genes have eukaryotic orthologs which are described in the present invention as targets to identify fungi and parasites. The eukaryotic homolog of elongation factor Tu is called elongation factor 1-alpha (EF-1a) (gene name: tef, tef1, ef1, ef-1 or EF-1). In fungi, the gene for EF-1 α occurs sometimes in two or more highly

similar duplicated copies (often named tef1, tef2, tef3...). In addition, eukaryotes have a copy of elongation factor Tu which is originating from their organelle genome ancestry (gene name: tuf1, tufM or tufA). For the purpose of the current invention, the genes for these four functionally and evolutionarily linked elongation factors (bacterial EF-Tu and EF-G, eukaryotic EF-1\alpha, and organellar EF-Tu) will hereafter be designated as **uf* nucleic acids and/or sequences**. The eukaryotic (mitochondrial) F0F1 type ATP-synthase beta subunit gene is named atp2 in yeast. For the purpose of the current invention, the genes of catalytic subunit of either F or V-type ATP-synthase will hereafter be designated as **atpD* nucleic acids and/or sequences**. The eukaryotic homologs of RecA are distributed in two families, typified by the Rad51 and Dmc1 proteins. Archaeal homologs of RecA are called RadA. For the purpose of the current invention, the genes corresponding to the latter proteins will hereafter be designated as **arecA* nucleic acids and/or sequences**.

In the description of this invention, the terms «nucleic acids» and «sequences» might be used interchangeably. However, «nucleic acids» are chemical entities while «sequences» are the pieces of information derived from (inherent to) these «nucleic acids». Both nucleic acids and sequences are equivalently valuable sources of information for the matter pertaining to this invention.

Analysis of multiple sequence alignments of tuf and atpD sequences permitted the design of oligonucleotide primers (and probes) capable of amplifying (or hybridizing to) segments of tuf (and/or fus) and atpD genes from a wide variety of bacterial species (see Examples 1 to 4, 24 and 26, and Table 7). Sequencing and amplification primer pairs for tuf nucleic acids and/or sequences are listed in Annex I and hybridization probes are listed in Annexes III and XLVII. Sequencing and amplification primer pairs for atpD nucleic acids and/or sequences are listed in Annex II. Analysis of the main subdivisions of tuf and atpD sequences (see Figures 1 and 2) permitted to design sequencing primers amplifying specifically each of these subdivisions. It should be noted that these sequencing primers could also be used as universal primers. However, since some of these sequencing primers

include several variable sequence (degenerated) positions, their sensitivity could be lower than that of universal primers developed for diagnostic purposes. Further subdivisions could be done on the basis of the various phyla where these genes are encountered.

Similarly, analysis of multiple sequence alignments of *recA* sequences present in the public databases permitted the design of oligonucleotide primers capable of amplifying segments of *recA* genes from a wide variety of bacterial species. Sequencing and amplification primer pairs for *recA* sequences are listed in Annex XXI. The main subdivisions of *recA* nucleic acids and/or sequences comprise *recA*, *rad51* and *dmc1*. Further subdivisions could be done on the basis of the various phyla where these genes are encountered.

The present inventor's strategy is to get as much sequence data information from the four conserved genes (tuf, fus, atpD and recA). This ensemble of sequence data forming a repertory (with subrepertories corresponding to each target gene and their main sequence subdivisions) and then using the sequence information of the sequence repertory (or subrepertories) to design primer pairs that could permit either universal detection of algae or archaea or bacteria or fungi or parasites, detection of a family or group of microorganism (e.g. Enterobacteriaceae), detection of a genus (e.g. Streptococcus) or finally a specific species (e.g. Staphylococcus aureus). It should be noted that for the purpose of the present invention a group of microorganisms is defined depending on the needs of the particular diagnostic test. It does not need to respect a particular taxonomical grouping or phylum. See Example 12 where primers were designed to amplify a group a bacteria consisting of the 17 major bacterial species encountered as contaminants of platelet concentrates. Also remark that in that Example, the primers are not only able to sensitively and rapidly detect at least the 17 important bacterial species, but could also detect other species as well, as shown in Table 14. In these circumstances the primers shown in Example 12 are considered universal for platelet-contaminating bacteria. To develop an assay specific for the latter, one or more primers or probes specific to each species could be designed. Another

example of primers and/or probes for group detection is given by the Pseudomonad group primers. These primers were designed based upon alignment of tuf sequences from real Pseudomonas species as well as from former Pseudomonas species such as Stenotrophomonas maltophilia. The resulting primers are able to amplify all Pseudomonas species tested as well as several species belonging to different genera, hence as being specific for a group including Pseudomonas and other species, we defined that group as Pseudomonads, as several members were former Pseudomonas.

For certain applications, it may be possible to develop a universal, group, family or genus-specific reaction and to proceed to species identification using sequence information within the amplicon to design species-specific internal probes or primers, or alternatively, to proceed directly by sequencing the amplicon. The various strategies will be discussed further below.

The ensembles formed by public and proprietary tuf, atpD and recA nucleic acids and/or sequences are used in a novel fashion so they constitute three databases containing useful information for the identification of microorganisms.

Sequence repertories of other gene targets were also built to solve some specific identification problems especially for microbial species genetically very similar to each other such as *E. coli* and *Shigella* (see Example 23). Based on *tuf*, *atpD* and *recA* sequences, *Streptococcus pneumoniae* is very difficult to differentiate from the closely related species *S. oralis* and *S. mitis*. Therefore, we elected to built a sequence repertory from *hexA* sequences (Example 19), a gene much more variable than our highly conserved *tuf*, *atpD* and *recA* nucleic acids and/or sequences.

For the detection of mutations associated with antibiotic resistance genes, we also built repertories to distinguish between point mutations reflecting only gene diversity and point mutations involved in resistance. This was done for *pbp1a*, *pbp2b* and *pbp2x* genes of penicillin-resistant and sensitive *Streptoccoccus pneumoniae* (Example 18) and also for *gyrA* and *parC* gene fragments of various bacterial species for which quinolone resistance is important to monitor.

Oligonucleotide primers and probes design and synthesis

The tuf, fus, atpD and recA DNA fragments sequenced by us and/or selected from public databases (GenBank and EMBL) were used to design oligonucleotides primers and probes for diagnostic purposes. Multiple sequence alignments were made using subsets of the tuf or atpD or recA sequences repertory. Subsets were chosen to encompass as much as possible of the targetted microorganism(s) DNA sequence data and also include sequence data from phylogenetically related microorganisms from which the targetted microorganism(s) should be distinguished. Regions suitable for primers and probes should be conserved for the targetted microorganism(s) and divergent for the microorganisms from which the targetted microorganism(s) should be distinguished. The large amount of tuf or atpD or recA sequences data in our repertory permits to reduce trial and errors in obtaining specific and ubiquitous primers and probes. We also relied on the corresponding peptide sequences of tuf, fus, atpD and recA nucleic acids and/or sequences to facilitate the identification of regions suitable for primers and probes design. As part of the design rules, all oligonucleotides (probes for hybridization and primers for DNA amplification by PCR) were evaluated for their suitability for hybridization or PCR amplification by computer analysis using standard programs (i.e. the Genetics Computer Group (GCG) programs and the primer analysis software OligoTM 5.0). The potential suitability of the PCR primer pairs was also evaluated prior to the synthesis by verifying the absence of unwanted features such as long stretches of one nucleotide and a high proportion of G or C residues at the 3' end (Persing et al., 1993, Diagnostic Molecular Microbiology: Principles and Society for Microbiology, Washington, Applications, American Oligonucleotide probes and amplification primers were synthesized using an automated DNA synthesizer (Perkin-Elmer Corp., Applied Biosystems Division).

The oligonucleotide sequence of primers or probes may be derived from either strand of the duplex DNA. The primers or probes may consist of the bases

A, G, C, or T or analogs and they may be degenerated at one or more chosen nucleotide position(s). The primers or probes may be of any suitable length and may be selected anywhere within the DNA sequences from proprietary fragments or from selected database sequences which are suitable for (i) the universal detection of algae or archaea or bacteria or fungi or parasites, (ii) the speciesspecific detection and identification of any microorganism, including but not limited to: Abiotrophia adiacens, Bacteroides fragilis, Bordetella pertussis, Candida dubliniensis. Candida glabrata, Candida Candida albicans. guilliermondii, Candida krusei, Candida lusitaniae, Candida parapsilosis, Candida tropicalis, Candida zeylanoides, Campylobacter jejuni and C. coli, Chlamydia pneumoniae, Chlamydia trachomatis, Cryptococcus neoformans, Cryptosporidium parvum, Enterococcus faecalis, Enterococcus faecium, Enterococcus gallinarum, Escherichia coli, Haemophilus influenzae, Legionella pneumophila, Mycoplasma pneumoniae, Neisseria gonorrhoeae, Pseudomonas aeruginosa, Staphylococcus aureus, Staphylococcus epidermidis, Staphylococcus hominis. saprophyticus, Staphylococcus Staphylococcus haemolyticus, Streptococcus agalactiae, Streptococcus pneumoniae, Trypanosoma brucei, Trypanosoma cruzi, (iii) the genus-specific detection of Bordetella species, Candida species, Clostridium species, Corynebacterium species, Cryptococcus species, Entamoeba species, Enterococcus species, Gemella species, Giardia species, Legionella species, Leishmania species, Staphylococcus species, Streptococcus species, Trypanosoma species, (iv) the family-specific detection of Enterobacteriaceae family members, Mycobacteriaceae family members, Trypanosomatidae family members, (v) the detection of Enterococcus Gemella and group, Enterococcus, casseliflavus-flavescens-gallinarum group, Pseudomonads extended group, Platelet-Abiotrophia adiacens contaminating bacteria group, (vi) the detection of clinically important antimicrobial agents resistance genes listed in Table 5, (vii) the detection of clinically important toxin genes listed in Table 6.

Variants for a given target microbial gene are naturally occurring and are attributable to sequence variation within that gene during evolution (Watson et al., 1987, Molecular Biology of the Gene, 4th ed., The Benjamin/Cummings Publishing Company, Menlo Park, CA; Lewin, 1989, Genes IV, John Wiley & Sons, New York, NY). For example, different strains of the same microbial species may have a single or more nucleotide variation(s) at the oligonucleotide hybridization site. The person skilled in the art is well aware of the existence of variant algal, archaeal, bacterial, fungal or parasitical DNA nucleic acids and/or sequences for a specific gene and that the frequency of sequence variations depends on the selective pressure during evolution on a given gene product. The detection of a variant sequence for a region between two PCR primers may be demonstrated by sequencing the amplification product. In order to show the presence of sequence variants at the primer hybridization site, one has to amplify a larger DNA target with PCR primers outside that hybridization site. Sequencing of this larger fragment will allow the detection of sequence variation at this site. A similar strategy may be applied to show variants at the hybridization site of a probe. Insofar as the divergence of the target nucleic acids and/or sequences or a part thereof does not affect the specificity and ubiquity of the amplification primers or probes, variant microbial DNA is under the scope of this invention. Variants of the selected primers or probes may also be used to amplify or hybridize to a variant DNA.

Sequencing of *tuf* nucleic acids and/or sequences from a variety of archaeal, bacterial, fungal and parasitical species

The nucleotide sequence of a portion of tuf nucleic acids and/or sequences was determined for a variety of archaeal, bacterial, fungal and parasitical species. The amplification primers (SEQ ID NOs. 664 and 697), which amplify a tuf gene portion of approximately 890 bp, were used along with newly designed sequencing primer pairs (See Annex I for the sequencing primers for tuf nucleic acids and/or

sequences). Most primer pairs can amplify different copies of tuf genes (tuf A and tufB). This is not surprising since it is known that for several bacterial species these two genes are nearly identical. For example, the entire tufA and tufB genes from E. coli differ at only 13 nucleotide positions (Neidhardtet al., 1996, Escherichia coli and Salmonella: Cellular and Molecular Biology, 2nd ed., American Society for Microbiology Press, Washington, D.C.). Similarly, some fungi are known to have two nearly identical copies of tuf nucleic acids and/or sequences (EF-1 α). These amplification primers are degenerated at several nucleotide positions and contain inosines in order to allow the amplification of a wide range of tuf nucleic acids and/or sequences. The strategy used to select these amplification primers is similar to that illustrated in Annex I for the selection of universal primers. The tuf sequencing primers even sometimes amplified highly divergent copies oftuf genes (tufC) as illustrated in the case of some enterococcal species (SEQ ID NOs.: 73, 75, 76, 614 to 618, 621 and 987 to 989). To prove this, we have determined the enterococcal tuf nucleic acids and/or sequences from PCR amplicons cloned into a plasmid vector. Using the sequence data from the cloned amplicons, we designed new sequencing primers specific to the divergent (tufC) copy of enterococci(SEQ ID NOs.: 658-659 and 661) and then sequenced directly the tufC amplicons. The amplification primers (SEQ ID NOs.: 543, 556, 557, 643-645, 660, 664, 694, 696 and 697) could be used to amplify the tuf nucleic acids and/or sequences from any bacterial species. The amplification primers (SEQ ID NOs.: 558, 559, 560, 653, 654, 655, 813, 815, 1974-1984, 1999-2003) could be used to amplify thetuf (EF-1α) genes from any fungal and/or parasitical species. The amplification primers SEQ ID NOs. 1221-1228 could be used to amplify bacterial tuf nucleic acids and/or sequences of the EF-G subdivision (fusA) (Figure 3). The amplification primers SEQ ID NOs. 1224, and 1227-1229 could be used to amplify bacterial tuf nucleic acids and/or sequences comprising the end of EF-G (fusA) and the beginning of EF-Tu (tuf), including the intergenic region, as shown in Figure 3. Most tuf fragments to be sequenced were amplified using the following amplification protocol: One µl of cell suspension (or of purified genomic DNA

 $0.1-100 \text{ ng/}\mu\text{l}$) was transferred directly to 19 μl of a PCR reaction mixture. Each PCR reaction contained 50 mM KCl, 10 mM Tris-HCl (pH 9.0), 0.1% Triton X-100, 2.5 mM MgCl₂, 1 μ M of each of the 2 primers, 200 μ M of each of the four dNTPs, 0.5 unit of Taq DNA polymerase (Promega Corp., Madison, WI). PCR reactions were subjected to cycling using a PTC-200 thermal cycler (MJ Research Inc., Watertown, Mass.) as follows: 3 min at 94-96 °C followed by 30-45 cycles of 1 min at 95 °C for the denaturation step, 1 min at 50-55 °C for the annealing step and 1 min at 72 °C for the extension step. Subsequently, twenty microliters of the PCR-amplified mixture were resolved by electrophoresis in a 1.5% agarose gel. The amplicons were then visualized by staining with methylene blue (Flores et al., 1992, Biotechniques, 13:203-205). The size of the amplification products was estimated by comparison with a 100-bp molecular weight ladder. The band corresponding to the specific amplification product was excised from the agarose gel and purified using the QIAquickTM gel extraction kit (QIAGEN Inc., Chatsworth, CA). The gel-purified DNA fragment was then used directly in the sequencing protocol. Both strands of the tuf genes amplification product were sequenced by the dideoxynucleotide chain termination sequencing method by using an Applied Biosystems automated DNA sequencer (model 377) with their Big DyeTM Terminator Cycle Sequencing Ready Reaction Kit (Applied Biosystems, Foster City, CA). The sequencing reactions were performed by using the same amplification primers and 10 ng/100 bp of the gel-purified amplicon per reaction. For the sequencing of long amplicons such as those of eukaryotic tuf (EF-1α) nucleic acids and/or sequences, we designed internal sequencing primers (SEQ ID NOs.: 654, 655 and 813) to be able to obtain sequence data on both strands for most of the fragment length. In order to ensure that the determined sequence did not contain errors attributable to the sequencing of PCR artefacts, we have sequenced two preparations of the gel-purified tuf amplification product originating from two independent PCR amplifications. For most target microbial species, the sequences determined for both amplicon preparations were identical. In case of discrepancies, amplicons from a third independent PCR amplification

were sequenced. Furthermore, the sequences of both strands were 100% complementary thereby confirming the high accuracy of the determined sequence. The *tuf* nucleic acids and/or sequences determined using the above strategy are described in the Sequence Listing. Table 7 gives the originating microbial species and the source for each *tuf* sequence in the Sequence Listing.

The alignment of the *tuf* sequences determined by us or selected from databases revealed clearly that the length of the sequenced portion of the *tuf* genes is variable. There may be insertions or deletions of several amino acids. In addition, in several fungi introns were observed. Intron nucleic acids and/or sequences are part of *tuf* nucleic acids and/or sequences and could be useful in the design of species-specific primers and probes. This explains why the size of the sequenced *tuf* amplification products was variable from one fungal species to another. Consequently, the nucleotide positions indicated on top of each of Annexes IV to XX, XXIII to XXXI, XXXVIII and XLII do not correspond for sequences having insertions or deletions.

It should also be noted that the various tuf nucleic acids and/or sequences determined by us occasionally contain base ambiguities. These degenerated nucleotides correspond to sequence variations between tufA and tufB genes (or copies of the EF-G subdivision of tuf nucleic acids and/or sequences, or copies of EF-1α subdivision of tuf nucleic acids and/or sequences for fungi and parasites) because the amplification primers amplify both tuf genes. These nucleotide variations were not attributable to nucleotide misincorporations by the Taq DNA polymerase because the sequence of both strands was identical and also because the sequences determined with both preparations of the gel-purified tuf amplicons obtained from two independent PCR amplifications were identical.

The selection of amplification primers from tuf nucleic acids and/or sequences

The tuf sequences determined by us or selected from public databases were used to select PCR primers for universal detection of bacteria, as well as for genus-

specific, species-specific family-specific or group-specific detection and identification. The strategy used to select these PCR primers was based on the analysis of multiple sequence alignments of various *tuf* sequences. For more details about the selection of PCR primers from *tuf* sequences please refer to Examples 5, 7-14, 17, 22, 24, 28, 30-31, 33, 36, and 38-40, and to Annexes VI-IX, XI-XIX and XXV.

Sequencing of atpD and recA nucleic acids and/or sequences from a variety of archaeal, bacterial, fungal and parasitical species

The method used to obtain atpD and recA nucleic acids and/or sequences is similar to that described above for tuf nucleic acids and/or sequences.

The selection of amplification primers from atpD or recA nucleic acids and/or sequences

The comparison of the nucleotide sequence for the *atpD* or *recA* genes from various archaeal, bacterial, fungal and parasitical species allowed the selection of PCR primers (refer to Examples 6, 13, 29, 34 and 37, and to Annexes IV, V, X, and XX).

DNA amplification

For DNA amplification by the widely used PCR (polymerase chain reaction) method, primer pairs were derived from proprietary DNA fragments or from database sequences. Prior to synthesis, the potential primer pairs were analyzed by using the OligoTM 5.0 software to verify that they were good candidates for PCR amplification.

During DNA amplification by PCR, two oligonucleotide primers binding respectively to each strand of the heat-denatured target DNA from the microbial

genome are used to amplify exponentially *in vitro* the target DNA by successive thermal cycles allowing denaturation of the DNA, annealing of the primers and synthesis of new targets at each cycle (Persing *et al*, 1993, Diagnostic Molecular Microbiology: Principles and Applications, American Society for Microbiology, Washington, D.C.).

Briefly, the PCR protocols were as follows: Treated clinical specimens or standardized bacterial or fungal or parasitical suspensions (see below) or purified genomic DNA from bacteria, fungi or parasites were amplified in a 20 μ 1 PCR reaction mixture. Each PCR reaction contained 50 mM KCl, 10 mM Tris-HCl (pH 9.0), 2.5 mM MgCl₂, 0.4 μ M of each primer, 200 μ M of each of the four dNTPs and 0.5 unit of Taq DNA polymerase (Promega) combined with the TaqStartTMantibody (Clontech Laboratories Inc., Palo Alto, CA). The TaqStartTM antibody, which is a neutralizing monoclonal antibody to Taq DNA polymerase, was added to all PCR reactions to enhance the specificity and the sensitivity of the amplifications (Kellogg et al., 1994, Biotechniques 16:1134-1137). The treatment of the clinical specimens varies with the type of specimen tested, since the composition and the sensitivity level required are different for each specimen type. It consists in a rapid protocol to lyse the microbial cells and eliminate or neutralize PCR inhibitors. For amplification from bacterial or fungal or parasitical cultures or from purified genomic DNA, the samples were added directly to the PCR amplification mixture without any pre-treatment step. An internal control was derived from sequences not found in the target microorganisms or in the human genome. The internal control was integrated into all amplification reactions to verify the efficiency of the PCR assays and to ensure that significant PCR inhibition was absent. Alternatively, an internal control derived from rRNA was also useful to monitor the efficiency of microbial lysis protocols.

PCR reactions were then subjected to thermal cycling (3 min at 94-96°C followed by 30 cycles of 1 second at 95°C for the denaturation step and 30 seconds at 50-65°C for the annealing-extension step) using a PTC-200 thermal cycler (MJ Research Inc.). The number of cycles performed for the PCR assays varies

according to the sensitivity level required. For example, the sensitivity level required for microbial detection directly from clinical specimens is higher for blood specimens than for urine specimens because the concentration of microorganisms associated with a septicemia can be much lower than that associated with a urinary tract infection. Consequently, more sensitive PCR assays having more thermal cycles are probably required for direct detection from blood specimens. Similarly, PCR assays performed directly from bacterial or fungal or parasitical cultures may be less sensitive than PCR assays performed directly from clinical specimens because the number of target organisms is normally much lower in clinical specimens than in microbial cultures.

The person skilled in the art of DNA amplification knows the existence of other rapid amplification procedures such as ligase chain reaction (LCR), transcription-mediated amplification (TMA), self-sustained sequence replication (3SR), nucleic acid sequence-based amplification (NASBA), strand displacement amplification (SDA), branched DNA (bDNA), cycling probe technology (CPT), solid phase amplification (SPA), rolling circle amplification technology (RCA), solid phase RCA, anchored SDA and nuclease dependent signal amplification (NDSA) (Lee et al., 1997, Nucleic Acid Amplification Technologies: Application to Disease Diagnosis, Eaton Publishing, Boston, MA; Persing et al., 1993, Diagnostic Molecular Microbiology: Principles and Applications, American Society for Microbiology, Washington, D.C.; Westin et al., 2000, Nat. Biotechnol. 18:199-204). The scope of this invention is not limited to the use of amplification by PCR, but rather includes the use of any rapid nucleic acid amplification method or any other procedure which may be used to increase the sensitivity and/or the rapidity of nucleic acid-based diagnostic tests. The scope of the present invention also covers the use of any nucleic acids amplification and detection technology including real-time or post-amplification detection technologies, any amplification technology combined with detection, any hybridization nucleic acid chips or arrays technologies, any amplification chips or combination of amplification and

hybridization chips technologies. Detection and identification by any sequencing method is also under the scope of the present invention.

Any oligonucleotide suitable for the amplification of nucleic acids by approaches other than PCR or for DNA hybridization which are derived from the species-specific, genus-specific and universal DNA fragments as well as from selected antimicrobial agents resistance or toxin gene sequences included in this document are also under the scope of this invention.

Detection of amplification products

Classically, detection of amplification is performed by standard ethidium bromide-stained agarose gel electrophoresis. It is clear that other methods for the detection of specific amplification products, which may be faster and more practical for routine diagnosis, may be used. Such methods may be based on the detection of fluorescence after or during amplification. One simple method for monitoring amplified DNA is to measure its rate of formation by measuring the increase in fluorescence of intercalating agents such as ethidium bromide or SYBR® Green I (Molecular Probes). If more specific detection is required, fluorescence-based technologies can monitor the appearance of a specific product during the reaction. The use of dual-labeled fluorogenic probes such as in the TaqMan[™] system (Applied Biosystems) which utilizes the 5'-3' exonuclease activity of the Taq polymerase is a good example (Livak K.J. et al. 1995, PCR Methods Appl. 4:357-362). TaqManTM can be performed during amplification and this "real-time" detection can be done in a single closed tube hence eliminating post-PCR sample handling and consequently preventing the risk of amplicon carryover. Several other fluorescence-based detection methods can be performed in real-time. Fluorescence resonance energy transfer (FRET) is the principle behind the use of adjacent hybridization probes (Wittwer, C.T. et al. 1997. BioTechniques 22:130-138), molecular beacons (Tyagi S. and Kramer F.R. 1996. Nature Biotechnology 14:303-308) and scorpions (Whitcomb et al. 1999. Nature

Biotechnology 17:804-807). Adjacent hybridization probes are designed to be internal to the amplification primers. The 3' end of one probe is labelled with a donor fluorophore while the 5' end of an adjacent probe is labelled with an acceptor fluorophore. When the two probes are specifically hybridized in closed proximity (spaced by 1 to 5 nucleotides) the donor fluorophore which has been excited by an external light source emits light that is absorbed by a second acceptor that emit more fluorescence and yields a FRET signal. Molecular beacons possess a stem-and-loop structure where the loop is the probe and at the bottom of the stem a fluorescent moiety is at one end while a quenching moiety is at the other end. The beacons undergo a fluorogenic conformational change when they hybridize to their targets hence separating the fluorochrome from its quencher. The FRET principle is also used in an air thermal cycler with a built-in fluorometer (Wittwer, C.T. et al. 1997. BioTechniques 22:130-138). The amplification and detection are extremely rapid as reactions are performed in capillaries: it takes only 18 min to complete 45 cycles. Those techniques are suitable especially in the case where few pathogens are searched for. Boehringer-Roche Inc. sells the LightCyclerTM, and Cepheid makes the SmartCycler. These two apparatus are capable of rapid cycle PCR combined with fluorescent SYBR® Green I or FRET detection. We recently demonstrated in our laboratory, real-time detection of 10 CFU in less than 40 minutes using adjacent hybridization probes on the LightCyclerTM. Methods based on the detection of fluorescence are particularly promising for utilization in routine diagnosis as they are very rapid, quantitative and can be automated.

Microbial pathogens detection and identification may also be performed by solid support or liquid hybridization using species-specific internal DNA probes hybridizing to an amplification product. Such probes may be generated from any sequence from our repertory and designed to specifically hybridize to DNA amplification products which are objects of the present invention. Alternatively, the internal probes for species or genus or family or group detection and identification may be derived from the amplicons produced by a universal, family-, group-, genus- or species-specific amplification assay(s). The oligonucleotide

probes may be labeled with biotin or with digoxigenin or with any other reporter molecule (for more details see below the section on hybrid capture). Hybrization on a solid support is amendable to miniaturization.

At present the oligonucleotide nucleic acid microarray technology is appealing. Currently, available low to medium density arrays (Heller et al., An integrated microelectronics hybridization system for genomic research and diagnostic applications. In: Harrison, D.J., and van den Berg, A., 1998, Micro total analysis systems '98, Kluwer Academic Publisher, Dordrecht.) could specifically capture fluorescent-labelled amplicons. Detection methods for hybridization are not limited to fluorescence; potentiometry, colorimetry and plasmon resonance are some examples of alternative detection methods. In addition to detection by hybridization, nucleic acid microarrays could be used to perform rapid sequencing by hybridization. Mass spectrometry could also be applicable for rapid identification of the amplicon or even for sequencing of the amplification products (Chiu and Cantor, 1999, Clinical Chemistry 45:1578; Berkenkamp et al., 1998, Science 281:260).

For the future of our assay format, we also consider the major challenge of molecular diagnostics tools, *i.e.*: integration of the major steps including sample preparation, genetic amplification, detection, data analysis and presentation (Anderson *et al.*, Advances in integrated genetic analysis. *In*: Harrison, D.J., and van den Berg, A., 1998, Micro total analysis systems 98, Kluwer Academic Publisher, Dordrecht.).

To ensure PCR efficiency, glycerol, dimethyl sulfoxide (DMSO) or other related solvents can be used to increase the sensitivity of the PCR and to overcome problems associated with the amplification of a target DNA having a high GC content or forming strong secondary structures (Dieffenbach and Dveksler, 1995, PCR Primer: A Laboratory Manual, Cold Spring Harbor Laboratory Press, Plainview, New York). The concentration ranges for glycerol and DMSO are 5-15% (v/v) and 3-10% (v/v), respectively. For the PCR reaction mixture, the concentration ranges for the amplification primers and MgCl₂ are 0.1-1.5 μ M and

1.0-10.0 mM, respectively. Modifications of the standard PCR protocol using external and nested primers (i.e. nested PCR) or using more than one primer pair (i.e. multiplex PCR) may also be used (Persing et al., 1993, Diagnostic Molecular Microbiology: Principles and Applications, American Society for Microbiology, Washington, D.C.). For more details about the PCR protocols and amplicon detection methods, see Examples.

Hybrid capture and chemiluminescence detection of amplification products

Hybridization and detection of amplicons by chemiluminescence were adapted from Nikiforov *et al.* (1994, PCR Methods and Applications 3:285-291 and 1995, Anal. Biochem. **227**:201-209) and from the DIGTM system protocol of Boehringer Mannheim. Briefly, 50 μl of a 25 picomoles solution of capture probe diluted in EDC {1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride} are immobilized in each well of 96-wells plates (MicroliteTM 2, Dynex) by incubation overnight at room temperature. The next day, the plates are incubated with a solution of 1% BSA diluted into TNTw (10 mM Tris-HCl, pH 7.5; 150 mM NaCl; 0.05% TweenTM 20) for 1 hour at 37 °C. The plates are then washed on a Wellwash AscentTM (Labsystems) with TNTw followed by Washing Buffer (100 mM maleic acid pH7.5; 150 mM NaCl; 0.3% TweenTM 20).

The amplicons were labelled with DIG-11-dUTP during PCR using the PCR DIG Labelling Mix from Boehringer Mannheim according to the manufacturer's instructions. Hybridization of the amplicons to the capture probes is performed in triplicate at stringent temperature (generally, probes are designed to allow hybrization at 55 °C, the stringent temperature) for 30 minutes in 1.5 M NaCl; 10 mM EDTA. It is followed by two washes in 2 X SSC; 0.1% SDS, then by four washes in 0.1X SSC; 0.1% SDS at the stringent temperature (55 °C). Detection with 1,2 dioxetane chemiluminescent alkaline phosphatase substrates like CSPD® (Tropix Inc.) is performed according to the manufacturer's instructions but with shorter incubations times and a different antibody concentration. The plates are

agitated at each step, the blocking incubation is performed for only 5 minutes, the anti-DIG-AP1 is used at a 1:1000 dilution, the incubation with antibody lasts 15 minutes, the plates are washed twice for only 5 minutes. Finally, after a 2 minutes incubation into the detection buffer, the plates are incubated 5 minutes with CSPD® at room temperature followed by a 10 minutes incubation at 37 °C without agitation. Luminous signal detection is performed on a Dynex Microtiter Plate Luminometer using RLU (Relative Light Units).

Specificity, ubiquity and sensitivity tests for oligonucleotide primers and probes

The specificity of oligonucleotide primers and probes was tested by amplification of DNA or by hybridization with bacterial or fungal or parasitical species selected from a panel comprising closely related species and species sharing the same anatomo-pathological site (see Annexes and Examples). All of the bacterial, fungal and parasitical species tested were likely to be pathogens associated with infections or potential contaminants which can be isolated from clinical specimens. Each target DNA could be released from microbial cells using standard chemical and/or physical treatments to lyse the cells (Sambrook *et al.*, 1989, Molecular Cloning: A Laboratory Manual, 2nd ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY) or alternatively, genomic DNA purified with the GNOMETM DNA kit (Bio101, Vista, CA) was used. Subsequently, the DNA was subjected to amplification with the primer pairs. Specific primers or probes amplified only the target microbial species, genus, family or group.

Oligonucleotides primers found to amplify specifically the target species, genus, family or group were subsequently tested for their ubiquity by amplification (i.e. ubiquitous primers amplified efficiently most or all isolates of the target species or genus or family or group). Finally, the sensitivity of the primers or probes was determined by using 10-fold or 2-fold dilutions of purified genomic DNA from the targeted microorganism. For most assays, sensitivity levels in the

range of 1-100 copies were obtained. The specificity, ubiquity and sensitivity of the PCR assays using the selected amplification primer pairs were tested either directly from cultures of microbial species or from purified microbial genomic DNA.

Probes were tested in hybrid capture assays as described above. An oligonucleotide probe was considered specific only when it hybridized solely to DNA from the species or genus or family or group from which it was selected. Oligonucleotide probes found to be specific were subsequently tested for their ubiquity (i.e. ubiquitous probes detected efficiently most or all isolates of the target species or genus or family or group) by hybridization to microbial DNAs from different clinical isolates of the species or genus or family or group of interest including ATCC reference strains. Similarly, oligonucleotide primers and probes could be derived from antimicrobial agents resistance or toxin genes which are objects of the present invention.

Reference strains

The reference strains used to build proprietary tuf, atpD and recA sequence data subrepertories, as well as to test the amplification and hybridization assays were obtained from (i) the American Type Culture Collection (ATCC), (ii) the Laboratoire de santé publique du Québec (LSPQ), (iii) the Centers for Disease Control and Prevention (CDC), (iv) the National Culture Type Collection (NCTC) and (v) several other reference laboratories throughout the world. The identity of our reference strains was confirmed by phenotypic testing and reconfirmed by analysis of tuf, atpD and recA sequences (see Example 13).

Antimicrobial agents resistance genes

Antimicrobial resistance complicates treatment and often leads to therapeutic failures. Furthermore, overuse of antibiotics inevitably leads to the emergence of

microbial resistance. Our goal is to provide clinicians, in approximately one hour, the needed information to prescribe optimal treatments. Besides the rapid identification of negative clinical specimens with DNA-based tests for universal algal, archaeal, bacterial, fungal or parasitical detection and the identification of the presence of a specific pathogen in the positive specimens with species- and/or genus- and/or family- and/or group-specific DNA-based tests, clinicians also need timely information about the ability of the microbial pathogen to resist antibiotic treatments. We feel that the most efficient strategy to evaluate rapidly microbial resistance to antimicrobials is to detect directly from the clinical specimens the most common and clinically important antimicrobial agents resistance genes (i.e. DNA-based tests for the specific detection of antimicrobial agents resistance genes). Since the sequence from the most important and common antimicrobial agents resistance genes are available from public databases, our strategy is to use the sequence from a portion or from the entire resistance gene to design specific oligonucleotide primers or probes which will be used as a basis for the development of sensitive and rapid DNA-based tests. The list of each of the antimicrobial agents resistance genes selected on the basis of their clinical relevance (i.e. high incidence and importance) is given in Table 5; descriptions of the designed amplification primers and internal probes are given in Annexes XXXIV-XXXVII, XXXIX, XLV, and L-LI. Our approach is unique because the antimicrobial agents resistance genes detection and the microbial detection and identification can be performed simultaneously, or independently, or sequentially in multiplex or parallel or sequential assays under uniform PCR amplification conditions. These amplifications can also be done separately.

Toxin genes

Toxin identification is often very important to prescribe optimal treatments. Besides the rapid identification of negative clinical specimens with DNA-based tests for universal bacterial detection and the identification of the presence of a

specific pathogen in the positive specimens with species- and/or genus- and/or family- and/or group-specific DNA-based tests, clinicians sometimes need timely information about the ability of certain bacterial pathogens to produce toxins. Since the sequence from the most important and common bacterial toxin genes are available from public databases, our strategy is to use the sequence from a portion or from the entire toxin gene to design specific oligonucleotide primers or probes which will be used as a basis for the development of sensitive and rapid DNAbased tests. The list of each of the bacterial toxin genes selected on the basis of their clinical relevance (i.e. high incidence and importance) is given in Table 6; descriptions of the designed amplification primers and internal probes are given in Annexes XXII, XXXII and XXXIII. Our approach is unique because the toxin genes detection and the bacterial detection and identification can be performed simultaneously, or independently, or sequentially, in multiplex or parallel or sequential assays under uniform PCR amplification conditions. These amplifications can also be done separately.

Universal bacterial detection

In the routine microbiology laboratory, a high percentage of clinical specimens sent for bacterial identification are negative by culture. Testing clinical samples with universal amplification primers or universal probes to detect the presence of bacteria prior to specific identification and screening out the numerous negative specimens is thus useful as it reduces costs and may rapidly orient the clinical management of the patients. Several amplification primers and probes were therefore synthesized from highly conserved portions of bacterial sequences from the *tuf*, *atpD* and *recA* nucleic acids and/or sequences. The universal primers selection was based on a multiple sequence alignment constructed with sequences from our repertory.

All computer analysis of amino acid and nucleotide sequences were performed by using the GCG programs. Subsequently, optimal PCR primers for

the universal amplification of bacteria were selected with the help of the OligoTM program. The selected primers are degenerated at several nucleotide positions and contain several inosines in order to allow the amplification of all clinically relevant bacterial species. Inosine is a nucleotide analog able to specifically bind to any of the four nucleotides A, C, G or T. Degenerated oligonucleotides consist of an oligonucleotide mix having two or more of the four nucleotides A, C, G or T at the site of mismatches. The inclusion of inosine and/or of base ambiguities in the amplification primers allow mismatch tolerance thereby permitting the amplification of a wider array of target nucleotide sequences (Dieffenbach and Dveksler, 1995 PCR Primer: A Laboratory Manual, Cold Spring Harbor Laboratory Press, Plainview, NY).

The amplification conditions with the universal primers are very similar to those used for the species- and genus-specific amplification assays except that the annealing temperature is slightly lower. The original universal PCR assay described in our assigned WO98/20157 (SEQ ID NOs. 23-24 of the latter application) was specific and nearly ubiquitous for the detection of bacteria. The specificity for bacteria was verified by amplifying genomic DNA isolated from the 12 fungal species as well as genomic DNA from Leishmania donovani, Saccharomyces cerevisiae and human lymphocytes. None of the above eukaryotic DNA preparations could be amplified by the universal assay, thereby suggesting that this test is specific for bacteria. The ubiquity of the universal assay was verified by amplifying genomic DNAs from 116 reference strains which represent 95 of the most clinically relevant bacterial species. These species have been selected from the bacterial species listed in Table 4. We found that at least 104 of these strains could be amplified. However, the assay could be improved since bacterial species which could not be amplified with the original tuf nucleic acids and/or sequences-based assay included species belonging to the following genera: Corynebacterium (11 species) and Stenotrophomonas (1 species). Sequencing of the tuf genes from these bacterial species and others has been performed in the scope of the present invention in order to improve the universal assay. This

sequencing data has been used to select new universal primers which may be more ubiquitous and more sensitive. Also, we improved our primer and probes design strategy by taking into consideration the phylogeny observed in analysing our repertory of tuf, atpD and recA sequences. Data from each of the 3 main subrepertories (tuf, atpD and recA) was subjected to a basic phylogenic analysis using the Pileup command from version 10 of the GCG package (Genetics Computer Group, inc.). This analysis indicated the main branches or phyla reflecting the relationships between sequences. Instead of trying to design primers or probes able to hybridize to all phyla, we designed primers or probes able to hybridize to the main phyla while trying to use the largest phylum possible. This strategy should allow less degenerated primers hence improving sensitivity and by combining primers in a mutiplex assay, improve ubiquity. Universal primers SEQ ID NOs. 643-645 based on tuf sequences have been designed to amplify most pathogenic bacteria except Actinomyceteae, Clostridiaceae and the Cytophaga, Flexibacter and Bacteroides phylum (pathogenic bacteria of this phylum include mostly Bacteroides, Porphyromonas and Prevotella species). Primers to fill these gaps have been designed for Actinomyceteae (SEQ ID NOs. 646-648), Clostridiaceae (SEQ ID NOs. 796-797, 808-811), and the Cytophaga, Flexibacter and Bacteroides phylum (SEQ ID NOs. 649-651), also derived from tuf nucleic acids and/or sequences. These primers sets could be used alone or in conjuction to render the universal assay more ubiquitous.

Universal primers derived from *atpD* sequences include SEQ ID NOs. 562-565. Combination of these primers does not amplify human DNA but should amplify almost all pathogenic bacterial species except proteobacteria belonging to the epsilon subdivision (*Campylobacter* and *Helicobacter*), the bacteria from the *Cytophaga*, *Flexibacter* and *Bacteroides* group and some actinomycetes and corynebacteria. By analysing *atpD* sequences from the latter species, primers and probes to specifically fill these gaps could be designed and used in conjuction with primers SEQ ID NOs. 562-565, also derived from *atpD* nucleic acids and/or sequences.

In addition, universality of the assay could be expanded by mixing atpD sequences-derived primers with tuf sequences-derived primers. Ultimately, even recA sequences-derived primers could be added to fill some gaps in the universal assay.

It is important to note that the 95 bacterial species selected to test the ubiquity of the universal assay include all of the most clinically relevant bacterial species associated with a variety of human infections acquired in the community or in hospitals (nosocomial infections). The most clinically important bacterial and fungal pathogens are listed in Tables 1 and 2.

Amino acid sequences derived from tuf, atpD and recA nucleic acids and/or sequences

The amino acid sequences translated from the repertory of tuf, atpD and recA nucleic acids and/or sequences are also an object of the present invention. The amino acid sequence data will be particularly useful for homology modeling of three-dimensional (3D) structure of the elongation factor Tu, elongation factor G, elongation factor 1a, ATPase subunit beta and RecA recombinase. For all these proteins, at least one structure model has been published using X-ray diffraction data from crystals. Based on those structural informations it is possible to use computer sofware to build 3D model structures for any other protein having peptide sequence homologies with the known structure (Greer, 1991, Methods in Enzymology, 202:239-252; Taylor, 1994, Trends Biotechnol., 12(5):154-158; Sali, 1995, Curr. Opin. Biotechnol. 6:437-451; Sanchez and Sali, 1997, Curr. Opin. Struct. Biol. 7:206-214; Fischer and Eisenberg, 1999, Curr. Opin. Struct. Biol. 9:208-211; Guex et al., 1999, Trends Biochem. Sci. 24: 364-367). Model structures of target proteins are used for the design or to predict the behavior of ligands and inhibitors such as antibiotics. Since EF-Tu and EF-G are already known as antibiotic targets (see above) and since the beta subunit of ATPase and RecA recombinase are essential to the survival of the microbial cells in natural

conditions of infection, all four proteins could be considered antibiotic targets. Sequence data, especially the new data generated by us could be very useful to assist the creation of new antibiotic molecules with desired spectrum of activity. In addition, model structures could be used to improve protein function for commercial purposes such as improving antibiotic production by microbial strains or increasing biomass.

The following detailed embodiments and appended drawings are provided as illustrative examples of his invention, with no intention to limit the scope thereof.

DESCRIPTION OF THE DRAWINGS

Figures 1 and 2 illustrate the principal subdivisions of the tuf and atpD sequences repertories, respectively. For the design of primers and probes, depending on the needs, one may want to use the complete data set illustrated on the top of the pyramid or use only a subset illustrated by the different branching points. Smaller subdivisions, representing groups, families, genus and species, could even be made to extend to the bottom of the pyramid. Because the tuf and atpD sequences are highly conserved and evolved with each species, the design of primers and probes does not need to include all the sequences within the database or its subdivisions. As illustrated in Annexes IV to XX, XXIII to XXXI, XXXVIII and XLII, depending on the use, sequences from a limited number of species can be carefully selected to represent: i) only the main phylogenetic branches from which the intended probes and primers need to be differentiating, and ii) only the species for which they need to be matching. However, for ubiquity purposes, and especially for primers and probes identifying large groups of species (genus, family, group or universal, or sequencing primers), the more data is included into the sequence analysis, the better the probes and primers will be suitable for each particular intended use. Similarly, for specificity purposes, a larger data set (or repertory) ensures optimal primers and probes design by reducing the chance of employing nonspecific oligonucleotides.

Figure 3 illustrates the approach used to design specific amplification primers from fusA as well as from the region between the end of fusA and the beginning of tuf in the streptomycin (str) operon (referred to as the fusA-tuf intergenic spacer in Table 7).

Figures 4 to 6 are illustrations to Example 42, whereas Figures 7 to 10 illustrate Example 43. Figures 11 and 12 illustrate Example 44.

FIGURE LEGENDS

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Figure 3. Schematic organization of universal amplification primers (SEQ ID NOs. 1221-1229) in the *str* operon. Amplicon sizes are given in bases pairs. Drawing not to scale, as the *fusA-tuf* intergenic spacer size varies depending on the bacterial species. Indicated amplicon lengths are for *E. coli*.

Figure 4. Abridged multiple amino acid sequence alignment of the partial tuf gene products from selected species illustrated using the program Alscript. Residues highly conserved in bacteria are boxed in grey and gaps are represented with dots. Residues in reverse print are unique to the enterococcal tufB as well as to streptococcal and lactococcal tuf gene products. Numbering is based on E. coli EF-Tu and secondary structure elements of E. coli EF-Tu are represented by cylinders (α -helices) and arrows (β -strands).

Figure 5. Distance matrix tree of bacterial EF-Tu based on amino acid sequence homology. The tree was constructed by the neighbor-joining method. The tree was rooted using archeal and eukaryotic EF- 1α genes as the outgroup. The scale bar represents 5% changes in amino acid sequence, as determined by taking the sum of all of the horizontal lines connecting two species.

Figure 6. Southern hybridization of BglII/XbaI digested genomic DNAs of some enterococci (except for E. casseliflavus and E. gallinarum whose genomic DNA was digested with BamHI/PvuII) using the tufA gene fragment of E. faecium as probes. The sizes of hybridizing fragments are shown in kilobases. Strains tested are listed in Table 16.

Figure 7. Pantoea and Tatumella species specific signature indel in atpD genes. The nucleotide positions given are for E. coli atpD sequence (GenBank accession no. V00267). Numbering starts from the first base of the initiation codon.

Figure 8: Trees based on sequence data from *tuf* (left side) and *atpD* (right side). The phylogenetic analysis was performed using the Neighbor-Joining method calculated using the Kimura two-parameter method. The value on each branch indicates the occurrence (%) of the branching order in 750 bootstrapped trees.

Figure 9: Phylogenetic tree of members of the family *Enterobacteriaceae* based on tuf (a), atpD (b), and 16S rDNA (c) genes. Trees were generated by neighborjoining method calculated using the Kimura two-parameter method. The value on each branch is the percentage of bootstrap replications supporting the branch. 750 bootstrap replications were calculated.

Figure 10: Plot of *tuf* distances versus 16S rDNA distances (a), atpD distances versus 16S rDNA distances (b), and atpD distances versus tuf distances (c). Symbols: O, distances between pairs of strains belonging to the same species; , distances between $E.\ coli$ strains and Shigella strains; \square , distances between pairs belonging to the same genus; \blacksquare , distances between pairs belonging to different genera; \triangle , distances between pairs belonging to different families.

EXAMPLES AND ANNEXES

For sake of clarity, here is a list of Examples and Annexes:

Example 1: Sequencing of bacterial atpD (F-type and V-type) gene fragments.

Example 2: Sequencing of eukaryotic atpD (F-type and V-type) gene fragments.

Example 3: Sequencing of eukaryotic *tuf* (EF-1) gene fragments.

Example 4: Sequencing of eukaryotic tuf (organelle origin, M) gene fragments.

- Example 5: Specific detection and identification of Streptococcus agalactiae using tuf sequences.
- Example 6: Specific detection and identification of Streptococcus agalactiae using atpD sequences.
- Example 7: Development of a PCR assay for detection and identification of staphylococci at genus and species levels.
- Example 8: Differentiating between the two closely related yeast species Candida albicans and Candida dubliniensis.
- Example 9: Specific detection and identification of Entamoeba histolytica.
- Example 10: Sensitive detection and identification of Chlamydia trachomatis.
- Example 11: Genus-specific detection and identification of enterococci.
- Example 12: Detection and identification of the major bacterial platelets contaminants using *tuf* sequences with a multiplex PCR test.
- Example 13: The resolving power of the *tuf* and *atpD* sequences databases is comparable to the biochemical methods for bacterial identification.
- Example 14: Detection of group B streptococci from clinical specimens.
- Example 15: Simultaneous detection and identification of Streptococcus pyogenes and its pyrogenic exotoxin A.
- Example 16: Real-time detection and identification of Shiga toxin-producing bacteria.
- Example 17: Development of a PCR assay for the detection and identification of staphylococci at genus and species levels and its associated *mecA* gene.
- Example 18: Sequencing of pbp1a, pbp2b and pbp2x genes of Streptoccoccus pneumoniae.
- Example 19: Sequencing of hexA genes of Streptococcus species.

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Example 20: Development of a multiplex PCR assay for the detection of Streptococcus pneumoniae and its penicillin resistance genes.

Example 21: Sequencing of the vancomycin resistance vanA, vanC1, vanC2 and vanC3 genes.

- Example 22: Development of a PCR assay for the detection and identification of enterococci at genus and species levels and its associated resistance genes *vanA* and *vanB*.
- Example 23: Development of a multiplex PCR assay for detection and identification of vancomycin-resistant Enterococcus faecalis, Enterococcus faecium, Enterococcus gallinarum, Enterococcus casseliflavus, and Enterococcus flavescens.
- Example 24: Universal amplification involving the EF-G (fusA) subdivision of tuf sequences.
- Example 25: DNA fragment isolation from *Staphylococcus saprophyticus* by arbitrarily primed PCR.
- Example 26: Sequencing of prokaryotic tuf gene fragments.
- Example 27: Sequencing of procaryotic recA gene fragments.
- Example 28: Specific detection and identification of *Escherichia coli/Shigella* sp. using *tuf* sequences.
- Example 29: Specific detection and identification of Klebsiella pneumoniae using atpD sequences.
- Example 30: Specific detection and identification of Acinetobacter baumanii using tuf sequences.
- Example 31: Specific detection and identification of Neisseria gonorrhoeae using tuf sequences.
- Example 32: Sequencing of bacterial gyrA and parC gene fragments.

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- Example 33: Development of a PCR assay for the specific detection and identification of *Staphylococcus aureus* and its quinolone resistance genes *gyrA* and *parC*.
- Example 34: Development of a PCR assay for the detection and identification of Klebsiella pneumoniae and its quinolone resistance genes gyrA and parC.

Example 35: Development of a PCR assay for the detection and identification of Streptococcus pneumoniae and its quinolone resistance genes gyrA and parC.

Example 36: Detection of extended-spectrum TEM-type β-lactamases in Escherichia coli.

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- Example 37: Detection of extended-spectrum SHV-type β-lactamases in Klebsiella pneumoniae.
- Example 38: Development of a PCR assay for the detection and identification of Neisseria gonorrhoeae and its associated tetracycline resistance gene tetM.
- Example 39: Development of a PCR assay for the detection and identification of Shigella sp. and their associated trimethoprim resistance gene dhfrla.
- Example 40: Development of a PCR assay for the detection and identification of Acinetobacter baumanii and its associated aminoglycoside resistance gene aph(3')-VIa.
- Example 41: Specific detection and identification of *Bacteroides fragilis* using atpD (V-type) sequences.
- Example 42: Evidence for horizontal gene transfer in the evolution of the elongation factor Tu in Enterococci.
- Example 43: Elongation factor Tu (tuf) and the F-ATPase beta-subunit (atpD) as phylogenetic tools for species of the family Enterobacteriaceae.
- Example 44: Testing new pairs of PCR primers selected from two species-specific genomic DNA fragments which are objects of US patent 6,001,564.
- Example 45: Testing modified versions of PCR primers derived from the sequence of several primers which are objects of US patent 6,001,564.

The various Annexes show the strategies used for the selection of a variety of DNA amplification primers, nucleic acid hybridization probes and molecular beacon internal probes:

(i) Annex I shows the amplification primers used for nucleic acid amplification from tuf sequences.

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- (ii) Annex II shows the amplification primers used for nucleic acid amplification from atpD sequences.
- (iii) Annex III shows the internal hybridization probes for detection of tuf sequences.
- (iv) Annex IV illustrates the strategy used for the selection of the amplification primers specific for atpD sequences of the F-type.
- (v) Annex V illustrates the strategy used for the selection of the amplification primers specific for *atpD* sequences of the V-type.
- (vi) Annex VI illustrates the strategy used for the selection of the amplification primers specific for the *tuf* sequences of organelle lineage (M, the letter M is used to indicate that in most cases, the organelle is the mitochondria).
- (vii) Annex VII illustrates the strategy used for the selection of the amplification primers specific for the tuf sequences of eukaryotes (EF-1).
- (viii) Annex VIII illustrates the strategy for the selection of *Streptococcus* agalactiae-specific amplification primers from tuf sequences.
- (ix) Annex IX illustrates the strategy for the selection of Streptococcus agalactiae-specific hybridization probes from tuf sequences.
- (x) Annex X illustrates the strategy for the selection of Streptococcus agalactiae-specific amplification primers from atpD sequences.
- (xi) Annex XI illustrates the strategy for the selection from tuf sequences of Candida albicans/dubliniensis-specific amplification primers, Candida albicans-specific hybridization probe and Candida dubliniensis-specific hybridization probe.

(xii) Annex XII illustrates the strategy for the selection of Staphylococcusspecific amplification primers from tuf sequences.

- (xiii) Annex XIII illustrates the strategy for the selection of the *Staphylococcus*-specific hybridization probe from *tuf* sequences.
- (xiv) Annex XIV illustrates the strategy for the selection of Staphylococcus saprophyticus-specific and Staphylococcus haemolyticus-specific hybridization probes from tuf sequences.
- (xv) Annex XV illustrates the strategy for the selection of Staphylococcus aureus-specific and Staphylococcus epidermidis-specific hybridization probes from tuf sequences.
- (xvi) Annex XVI illustrates the strategy for the selection of the Staphylococcus hominis-specific hybridization probe from tuf sequences.
- (xvii) Annex XVII illustrates the strategy for the selection of the *Enterococcus*-specific amplification primers from *tuf* sequences.
- (xviii) Annex XVIII illustrates the strategy for the selection of the Enterococcus faecalis-specific hybridization probe, of the Enterococcus faecium-specific hybridization probe and of the Enterococcus casseliflavus-flavescens-gallinarum group-specific hybridization probe from tuf sequences.
- (xix) Annex XIX illustrates the strategy for the selection of primers from tuf sequences for the identification of platelets contaminants.
- (xx) Annex XX illustrates the strategy for the selection of the universal amplification primers from atpD sequences.
- (xxi) Annex XXI shows the amplification primers used for nucleic acid amplification from recA sequences.
- (xxii) Annex XXII shows the specific and ubiquitous primers for nucleic acid amplification from speA sequences.
- (xxiii) Annex XXIII illustrates the first strategy for the selection of Streptococcus pyogenes-specific amplification primers from speA sequences.

(xxiv) Annex XXIV illustrates the second strategy for the selection of Streptococcus pyogenes-specific amplification primers from speA sequences.

(xxv) Annex XXV illustrates the strategy for the selection of *Streptococcus* pyogenes-specific amplification primers from tuf sequences.

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- (xxvi) Annex XXVI illustrates the strategy for the selection of stx_1 -specific amplification primers and hybridization probe.
- (xxvii) Annex XXVII illustrates the strategy for the selection of stx_2 -specific amplification primers and hybridization probe.
- (xxviii) Annex XXVIII illustrates the strategy for the selection of vanA-specific amplification primers from van sequences.
- (xxix) Annex XXIX illustrates the strategy for the selection of vanB-specific amplification primers from van sequences.
- (xxx) Annex XXX illustrates the strategy for the selection of vanC-specific amplification primers from vanC sequences.
- (xxxi) Annex XXXI illustrates the strategy for the selection of *Streptococcus* pneumoniae-specific amplification primers and hybridization probes from pbpla sequences.
- (xxxii) Annex XXXII shows the specific and ubiquitous primers for nucleic acid amplification from toxin gene sequences.
- (xxxiii) Annex XXXIII shows the molecular beacon internal hybridization probes for specific detection of toxin sequences.
- (xxxiv) Annex XXXIV shows the specific and ubiquitous primers for nucleic acid amplification from van sequences.
- (xxxv) Annex XXXV shows the internal hybridization probes for specific detection of van sequences.
- (xxxvi) Annex XXXVI shows the specific and ubiquitous primers for nucleic acid amplification from pbp sequences.
- (xxxvii) Annex XXXVII shows the internal hybridization probes for specific detection of pbp sequences.

(xxxviii)Annex XXXVIII illustrates the strategy for the selection of vanAB-specific amplification primers and vanA- and vanB- specific hybridization probes from van sequences.

(xxxix) Annex XXXIX shows the internal hybridization probe for specific detection of mecA.

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- (xl) Annex XL shows the specific and ubiquitous primers for nucleic acid amplification from hexA sequences.
- (xli) Annex XLI shows the internal hybridization probe for specific detection of hexA.
- (xlii) Annex XLII illustrates the strategy for the selection of *Streptococcus* pneumoniae species-specific amplification primers and hybridization probe from hexA sequences.
- (xliii) Annex XLIII shows the specific and ubiquitous primers for nucleic acid amplification from pcp sequences.
- (xliv) Annex XLIV shows specific and ubiquitous primers for nucleic acid amplification of S. saprophyticus sequences of unknown coding potential.
- (xlv) Annex XLV shows the molecular beacon internal hybridization probes for specific detection of antimicrobial agents resistance gene sequences.
- (xlvi) Annex XLVI shows the molecular beacon internal hybridization probe for specific detection of S. aureus gene sequences of unknown coding potential.
- (xlvii) Annex XLVII shows the molecular beacon hybridization internal probe for specific detection of *tuf* sequences.
- (xlviii) Annex XLVIII shows the molecular beacon internal hybridization probes for specific detection of *ddl* and *mtl* sequences.
- (xlix) Annex XLIX shows the internal hybridization probe for specific detection of S. aureus sequences of unknown coding potential.
- (l) Annex L shows the amplification primers used for nucleic acid amplification from antimicrobial agents resistance genes sequences.

(li) Annex LI shows the internal hybridization probes for specific detection of antimicrobial agents resistance genes sequences.

- (lii) Annex LII shows the molecular beacon internal hybridization probes for specific detection of *atpD* sequences.
- (liii) Annex LIII shows the internal hybridization probes for specific detection of atpD sequences.
- (liv) Annex LIVI shows the internal hybridization probes for specific detection of ddl and mtl sequences.

As shown in these Annexes, the selected amplification primers may contain inosines and/or base ambiguities. Inosine is a nucleotide analog able to specifically bind to any of the four nucleotides A, C, G or T. Alternatively, degenerated oligonucleotides which consist of an oligonucleotide mix having two or more of the four nucleotides A, C, G or T at the site of mismatches were used. The inclusion of inosine and/or of degeneracies in the amplification primers allows mismatch tolerance thereby permitting the amplification of a wider array of target nucleotide sequences (Dieffenbach and Dveksler, 1995 PCR Primer: A Laboratory Manual, Cold Spring Harbor Laboratory Press, Plainview, New York).

EXAMPLES

EXAMPLE 1:

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Sequencing of bacterial atpD (F-type and V-type) gene fragments. As shown in Annex IV, the comparison of publicly available atpD (F-type) sequences from a variety of bacterial species revealed conserved regions allowing the design of PCR primers able to amplify atpD sequences (F-type) from a wide range of bacterial species. Using primers pairs SEQ ID NOs. 566 and 567, 566 and 814, 568 and 567, 570 and 567, 572 and 567, 569 and 567, 571 and 567, 700 and 567, it was possible to amplify and sequence atpD sequences SEQ ID NOs. 242-270, 272-398, 673-

674, 737-767, 866-867, 942-955, 1245-1254, 1256-1265, 1527, 1576, 1577, 1600-1604, 1640-1646, 1649, 1652, 1655, 1657, 1659-1660, 1671, 1844-1845, and 1849-1865.

Similarly, Annex V shows the strategy to design the PCR primers able to amplify atpD sequences of the V-type from a wide range of archaeal and bacterial species. Using primers SEQ ID NOs. 681-683, it was possible to amplify and sequence atpD sequences SEQ ID NOs. 827-832, 929-931, 958 and 966. As the gene was difficult to amplify for several species, additional amplification primers were designed inside the original amplicon (SEQ ID NOs. 1203-1207) in order to obtain sequence information for these species. Other primers (SEQ ID NO. 1212, 1213, 2282-2285) were also designed to amplify regions of the atpD gene (V-type) in archaebacteria.

EXAMPLE 2:

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Sequencing of eukaryotic atpD (F-type and V-type) gene fragments. The comparison of publicly available atpD (F-type) sequences from a variety of fungal and parasitical species revealed conserved regions allowing the design of PCR primers able to amplify atpD sequences from a wide range of fungal and parasitical species. Using primers pairs SEQ ID NOs. 568 and 573, 574 and 573, 574 and 708, and 566 and 567, it was possible to amplify and sequence atpD sequences SEQ ID NOs. 458-497, 530-538, 663, 667, 676, 678-680, 768-778, 856-862, 889-896, 941, 1638-1639, 1647, 1650-1651, 1653-1654, 1656, 1658, 1684, 1846-1848, and 2189-2192.

In the same manner, the primers described in Annex V (SEQ ID NOs. 681-683) could amplify the *atpD* (V-type) gene from various fungal and parasitical species. This strategy allowed to obtain SEQ ID NOs. 834-839, 956-957, and 959-965.

WO 01/23604 EXAMPLE 3:

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Sequencing of eukaryotic *tuf* (EF-1) gene fragments. As shown in Annex VII, the comparison of publicly available *tuf* (EF-1) sequences from a variety of fungal and parasitical species revealed conserved regions allowing the design of PCR primers able to amplify *tuf* sequences from a wide range of fungal and parasitical species. Using primers pairs SEQ ID NOs. 558 and 559, 813 and 559, 558 and 815, 560 and 559, 653 and 559, 558 and 655, and 654 and 559, 1999 and 2000, 2001 and 2003, 2002 and 2003, it was possible to amplify and sequence *tuf* sequences SEQ ID NOs. 399-457, 509-529, 622-624, 677, 779-790, 840-842, 865, 897-903, 1266-1287, 1561-1571 and 1685.

EXAMPLE 4:

Sequencing of eukaryotic *tuf* (organelle origin, M) gene fragments. As shown in Annex VI, the comparison of publicly available *tuf* (organelle origin, M) sequences from a variety of fungal and parasitical organelles revealed conserved regions allowing the design of PCR primers able to amplify *tuf* sequences of several organelles belonging to a wide range fungal and parasitical species. Using primers pairs SEQ ID NOs. 664 and 652, 664 and 561, 911 and 914, 912 and 914, 913 and 915, 916 and 561, 664 and 917, it was possible to amplify and sequence *tuf* sequences SEQ ID NOs. 498-508, 791-792, 843-855, 904-910, 1664, 1666-1667, 1669-1670, 1673-1683, 1686-1689, 1874-1876, 1879, 1956-1960, and 2193-2199.

EXAMPLE 5:

Specific detection and identification of Streptococcus agalactiae using tuf sequences. As shown in Annex VIII, the comparison of tuf sequences from a variety of bacterial species allowed the selection of PCR primers specific for S. agalactiae. The strategy used to design the PCR primers was based on the analysis

of a multiple sequence alignment of various tuf sequences. The multiple sequence alignment includes the tuf sequences of four bacterial strains from the target species as well as tuf sequences from other species and bacterial genera, especially representatives of closely related species. A careful analysis of this alignment allowed the selection of oligonucleotide sequences which are conserved within the target species but which discriminate sequences from other species and genera, especially from the closely related species, thereby permitting the species-specific, ubiquitous and sensitive detection and identification of the target bacterial species.

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The chosen primer pair, oligos SEQ ID NO. 549 and SEQ ID NO. 550, gives an amplification product of 252 bp. Standard PCR was carried out using 0.4 μ M of each primer, 2.5 mM MgCl₂, BSA 0.05 mM, 1X Taq Buffer (Promega), dNTP 0.2 mM (Pharmacia), 0,5 U *Taq* DNA polymerase (Promega) coupled with TaqStartTM antibody (Clontech Laboratories Inc., Palo Alto), 1 μ l of genomic DNA sample in a final volume of 20 μ l using a PTC-200 thermocycler (MJ Research Inc.). The optimal cycling conditions for maximum sensitivity and specificity were 3 minutes at 95 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 62 °C, followed by terminal extension at 72 °C for 2 minutes. Detection of the PCR products was made by electrophoresis in agarose gels (2 %) containing 0.25 μ g/ml of ethidium bromide.

Specificity of the assay was tested by adding into the PCR reactions, 0.1 ng of genomic DNA from each of the bacterial species listed in Table 8. Efficient amplification was observed only for the 5 S. agalactiae strains listed. Of the other bacterial species, including 32 species representative of the vaginal flora and 27 other streptococcal species, only S. acidominimus yielded amplification. The signal with 0.1 ng of S. acidominimus genomic DNA was weak and the detection limit for this species was 10 pg (corresponding to more than 4000 genome copies) while the detection limit for S. agalactiae was 2.5 fg (corresponding to one genome copy) of genomic DNA.

To increase the specificity of the assay, internal probes were designed for FRET (Fluorescence Resonance Energy Transfer) detection using the LightCycler™ (Idaho Technology). As illustrated in Annex IX, a multiple sequence alignment of streptococcal *tuf* sequence fragments corresponding to the 252 bp region amplified by primers SEQ ID NO. 549 and SEQ ID NO. 550, was used for the design of internal probes TSagHF436 (SEQ ID NO. 582) and TSagHF465 (SEQ ID NO. 583). The region of the amplicon selected for internal probes contained sequences unique and specific to *S. agalactiae*. SEQ ID NO. 583, the more specific probe, is labelled with fluorescein in 3', while SEQ ID NO. 582, the less discriminant probe, is labelled with CY5 in 5' and blocked in 3' with a phosphate group. However, since the FRET signal is only emitted if both probes are adjacently hybridized on the same target amplicon, detection is highly specific.

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Real-time detection of PCR products using the LightCyclerTM was carried out using 0.4 μ M of each primer (SEQ ID NO. 549-550), 0.2 μ M of each probe (SEQ ID NO. 582-583), 2.5 mM MgCl₂, BSA 450 μ g/ml, 1X PC2 Buffer (AB Peptides, St-Louis, MO), dNTP 0.2 mM (Pharmacia), 0.5 U KlenTaq1™ DNA polymerase (AB Peptides) coupled with TaqStartTM antibody (Clontech Laboratories Inc., Palo Alto), 0.7 μ l of genomic DNA sample in a final volume of 7 μ l using a LightCycler thermocycler (Idaho Technology). The optimal cycling conditions for maximum sensitivity and specificity were 3 minutes at 94 °C for initial denaturation, then forty cycles of three steps consisting of 0 second (this setting meaning the LightCycler will reach the target temperature and stay at it for its minimal amount of time) at 94 °C, 10 seconds at 64 °C, 20 seconds at 72 °C. Amplification was monitored during each annealing steps using the fluorescence ratio. The streptococcal species having close sequence homologies with the tuf sequence of S. agalactiae (S. acidominimus, S. anginosus, S. bovis, S. dysgalactiae, S. equi, S. ferus, S. gordonii, S. intermedius, S. parasanguis, S. parauberis, S. salivarius, S. sanguis, S. suis) as well as S. agalactiae were tested in the

LightCycler with 0.07 ng of genomic DNA per reaction. Only *S. agalactiae* yielded an amplification signal, hence demonstrating that the assay is species-specific. With the LightCyclerTM assay using the internal FRET probes, the detection limit for *S. agalactiae* was 1-2 genome copies of genomic DNA.

EXAMPLE 6:

Specific detection and identification of Streptococcus agalactiae using atpD sequences. As shown in Annex X, the comparison of atpD sequences from a variety of bacterial species allowed the selection of PCR primers specific for S. agalactiae. The primer design strategy is similar to the strategy described in the preceding Example except that atpD sequences were used in the alignment.

Four primers were selected, ASag42 (SEQ ID NO. 627), ASag52 (SEQ ID NO. 628), ASag206 (SEQ ID NO. 625) and ASag371 (SEQ ID NO. 626). The following combinations of these four primers give four amplicons; SEQ ID NO. 627 + SEQ ID NO. 625 = 190 bp, SEQ ID NO. 628 + SEQ ID NO. 625 = 180 bp, SEQ ID NO. 627 + SEQ ID NO. 626 = 355 bp, and SEQ ID NO. 628 + SEQ ID NO. 626 = 345 bp.

Standard PCR was carried out on PTC-200 thermocyclers (MJ Research Inc) using 0.4 μ M of each primers pair, 2.5 mM MgCl₂, BSA 0.05 mM, 1X taq Buffer (Promega), dNTP 0.2 mM (Pharmacia), 0.5 U Taq DNA polymerase (Promega) coupled with TaqStartTM antibody (Clontech Laboratories Inc., Palo Alto), 1 μ l of genomic DNA sample in a final volume of 20 μ L. The optimal cycling conditions for maximum sensitivity and specificity were adjusted for each primer pair. Three minutes at 95 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at the optimal annealing temperature specified below were followed by terminal extension at 72 °C for 2 minutes. Detection of the PCR products was made by electrophoresis in agarose gels (2 %) containing

 $0.25 \mu g/ml$ of ethidium bromide. Since atpD sequences are relatively more specific than tuf sequences, only the most closely related species namely, the steptococcal species listed in Table 9, were tested.

All four primer pairs only amplified the six *S. agalactiae* strains. With an annealing temperature of 63 °C, the primer pair SEQ ID NO. 627 + SEQ ID NO. 625 had a sensitivity of 1-5 fg (equivalent to 1-2 genome copies). At 55 °C, the primer pair SEQ ID NO. 628 + SEQ ID NO. 625 had a sensitivity of 2.5 fg (equivalent to 1 genome copy). At 60 °C, the primer pair SEQ ID NO. 627 + SEQ ID NO. 626 had a sensitivity of 10 fg (equivalent to 4 genome copies). At 58 °C, the primer pair SEQ ID NO. 628 + SEQ ID NO. 626 had a sensitivity of 2.5-5 fg (equivalent to 1-2 genome copies). This proves that all four primer pairs can detect *S. agalactiae* with high specificity and sensitivity. Together with Example 5, this example demonstrates that both *tuf* and *atpD* sequences are suitable and flexible targets for the identification of microorganisms at the species level. The fact that 4 different primer pairs based on *atpD* sequences led to efficient and specific amplification of *S. agalactiae* demonstrates that the challenge is to find target genes suitable for diagnostic purposes, rather than finding primer pairs from these target sequences.

EXAMPLE 7:

Development of a PCR assay for detection and identification of staphylococci at genus and species levels.

Materials and Methods

Bacterial strains. The specificity of the PCR assay was verified by using a panel of ATCC (America Type Culture Collection) and DSMZ (Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH; German Collection of

Microorganisms and Cell Cultures) reference strains consisting of 33 gramnegative and 47 gram-positive bacterial species (Table 12). In addition, 295 clinical isolates representing 11 different species of staphylococci from the microbiology laboratory of the Centre Hospitalier Universitaire de Québec, Pavillon Centre Hospitalier de l'Université Laval (CHUL) (Ste-Foy, Québec, Canada) were also tested to further validate the *Staphylococcus*-specific PCR assay. These strains were all identified by using (i) conventional methods or (ii) the automated MicroScan Autoscan-4 system equipped with the Positive BP Combo Panel Type 6 (Dade Diagnostics, Mississauga, Ontario, Canada). Bacterial strains from frozen stocks kept at -80 °C in brain heart infusion (BHI) broth containing 10% glycerol were cultured on sheep blood agar or in BHI broth (Quelab Laboratories Inc, Montréal, Québec, Canada).

PCR primers and internal probes. Based on multiple sequence alignments, regions of the *tuf* gene unique to staphylococci were identified. *Staphylococcus*-specific PCR primers TStaG422 (SEQ ID NO. 553) and TStaG765 (SEQ ID NO. 575) were derived from these regions (Annex XII). These PCR primers are displaced by two nucleotide positions compared to original *Staphylococcus*-specific PCR primers described in our patent publication WO98/20157 (SEQ ID NOs. 17 and 20 in the said patent publication). These modifications were done to ensure specificity and ubiquity of the primer pair, in the light of new *tuf* sequence data revealed in the present patent application for several additional staphylococcal species and strains.

Similarly, sequence alignment analysis were performed to design genus and species-specific internal probes (see Annexes XIII to XVI). Two internal probes specific for *Staphylococcus* (SEQ ID NOs. 605-606), five specific for *S. aureus* (SEQ ID NOs. 584-588), five specific for *S. epidermidis* (SEQ ID NO. 589-593), two specific for *S. haemolyticus* (SEQ ID NOs. 594-595), three specific for *S. hominis* (SEQ ID NOs. 596-598), four specific for *S. saprophyticus* (SEQ ID NOs. 599-601 and 695), and two specific for coagulase-negative *Staphylococcus* species including

S. epidermidis, S. hominis, S. saprophyticus, S. auricularis, S. capitis, S. haemolyticus, S. lugdunensis, S. simulans, S. cohnii and S. warneri (SEQ ID NOs. 1175-1176) were designed. The range of mismatches between the Staphylococcusspecific 371-bp amplicon and each of the 20-mer species-specific internal probes was from 1 to 5, in the middle of the probe when possible. No mismatches were present in the two Staphylococcus-specific probes for the 11 species analyzed: S. aureus, S. auricularis, S. capitis, S. cohnii, S. epidermidis, S. haemolyticus, S. hominis, S. lugdunensis, S. saprophyticus, S. simulans and S. warneri. In order to verify the intra-specific sequence conservation of the nucleotide sequence, sequences were obtained for the 371-bp amplicon from five unrelated ATCC and clinical strains for each of the species S. aureus, S. epidermidis, S. haemolyticus, S. hominis and S. saprophyticus. The OligoTM (version 5.0) primer analysis software (National Biosciences, Plymouth, Minn.) was used to confirm the absence of selfcomplementary regions within and between the primers or probes. When required, the primers contained inosines or degenerated nucleotides at one or more variable positions. Oligonucleotide primers and probes were synthesized on a model 394 DNA synthesizer (Applied Biosystems, Mississauga, Ontario, Canada). Detection of the hybridization was performed with the DIG-labeled dUTP incorporated during amplification with the Staphylococcus-specific PCR assay, and the hybridization signal was detected with a luminometer (Dynex Technologies) as described above in the section on luminescent detection of amplification products. Annexes XIII to XVI illustrate the strategy for the selection of several internal probes.

PCR amplification. For all bacterial species, amplification was performed from purified genomic DNA or from a bacterial suspension whose turbidity was adjusted to that of a 0.5 McFarland standard, which corresponds to approximately 1.5 x 10⁸ bacteria per ml. One nanogram of genomic DNA or 1 µl of the standardized bacterial suspension was transferred directly to a 19 µl PCR mixture. Each PCR reaction contained 50 mM KCl, 10 mM Tris-HCl (pH 9.0), 0.1% Triton X-100, 2.5 mM

MgCl₂, 0.2 μ M (each) of the two *Staphylococcus* genus-specific primers (SEQ ID NOs. 553 and 575), 200 μ M (each) of the four deoxynucleoside triphosphates (Pharmacia Biotech), 3.3 μ g/ μ l bovine serum albumin (BSA) (Sigma-Aldrich Canada Ltd, Oakville, Ontario, Canada), and 0.5 U *Taq* polymerase (Promega) coupled with *Taq*StartTM Antibody (Clontech). The PCR amplification was performed as follows: 3 min. at 94 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 55 °C, plus a terminal extension at 72 °C for 2 minutes. Detection of the PCR products was made by electrophoresis in agarose gels (2 %) containing 0.25 μ g/ml of ethidium bromide. Visualization of the PCR products was made under UV at 254 nm.

For determination of the sensitivities of the PCR assays, two-fold dilutions of purified genomic DNA were used to determine the minimal number of genome copies which can be detected.

Results

Amplifications with the Staphylococcus genus-specific PCR assay. The specificity of the assay was assessed by performing 30-cycle and 40-cycle PCR amplifications with the panel of gram-positive (47 species from 8 genera) and gram-negative (33 species from 22 genera) bacterial species listed in Table 12. The PCR assay was able to detect efficiently 27 of 27 staphylococcal species tested in both 30-cycle and 40-cycle regimens. For 30-cycle PCR, all bacterial species tested other than staphylococci were negative. For 40-cycle PCR, Enterococcus faecalis and Macrococcus caseolyticus were slightly positive for the Staphylococcus-specific PCR assay. The other species tested remained negative. Ubiquity tests performed on a collection of 295 clinical isolates provided by the microbiology laboratory of the Centre Hospitalier Universitaire de Québec, Pavillon Centre Hospitalier de l'Université Laval (CHUL), including Staphylococcus aureus (n=34), S. auricularis (n=2), S. capitis (n=19), S. cohnii (n=5), S. epidermidis (n=18), S. haemolyticus

(n=21), S. hominis (n=73), S. lugdunensis (n=17), S. saprophyticus (n=6), S. simulans (n=3), S. warneri (n=32) and Staphylococcus sp. (n=65), showed a uniform amplification signal with the 30-cycle PCR assays and a perfect relation between the genotype and classical identification schemes.

The sensitivity of the *Staphylococcus*-specific assay with 30-cycle and 40-cycle PCR protocols was determined by using purified genomic DNA from the 11 staphylococcal species previously mentioned. For PCR with 30 cycles, a detection limit of 50 copies of genomic DNA was consistently obtained. In order to enhance the sensitivity of the assay, the number of cycles was increased. For 40-cycle PCR assays, the detection limit was lowered to a range of 5-10 genome copies, depending on the staphylococcal species tested.

Hybridization between the Staphylococcus-specific 371-bp amplicon and species-specific or genus-specific internal probes. Inter-species polymorphism was sufficient to generate species-specific internal probes for each of the principal species involved in human diseases (S. aureus, S. epidermidis, S. haemolyticus, S. hominis and S. saprophyticus). In order to verify the intra-species sequence conservation of the nucleotide sequence, sequence comparisons were performed on the 371-bp amplicon from five unrelated ATCC and clinical strains for each of the 5 principal staphylococcal species: S. aureus, S. epidermidis, S. haemolyticus, S. hominis and S. saprophyticus. Results showed a high level of conservation of nucleotide sequence between different unrelated strains from the same species. This sequence information allowed the development of staphylococcal species identification assays using species-specific internal probes hybridizing to the 371bp amplicon. These assays are specific and ubiquitous for those five staphylococcal species. In addition to the species-specific internal probes, the genus-specific internals probes were able to recognize all or most Staphylococcus species tested.

EXAMPLE 8:

Differentiating between the two closely related yeast species Candida albicans and Candida dubliniensis. It is often useful for the clinician to be able to differentiate between two very closely related species of microorganisms. Candida albicans is the most important cause of invasive human mycose. In recent years, a very closely related species, Candida dubliniensis, was isolated in immunosuppressed patients. These two species are difficult to distinguish by classic biochemical methods. This example demonstrates the use of tuf sequences to differentiate Candida albicans and Candida dubliniensis. PCR primers SEQ ID NOs. 11-12, from previous patent publication WO98/20157, were selected for their ability to specifically amplify a tuf (elongation factor 1 alpha type) fragment from both species (see Annex XI for primer positions). Within this tuf fragment, a region differentiating C. albicans and C. dubliniensis by two nucleotides was selected and used to design two internal probes (see Annex XI for probe design, SEQ ID NOs. 577 and 578) specific for each species. Amplification of genomic DNA from C. albicans and C. dubliniensis was carried out using DIG-11-dUTP as described above in the section on chemiluminescent detection of amplification products. Internal probes SEQ ID NOs. 577 and 578 were immobilized on the bottom of individual microtiter plates and hybridization was carried out as described above in the above section on chemiluminescent detection of amplification products. Luminometer data showed that the amplicon from C. albicans hybridized only to probe SEQ ID NO. 577 while the amplicon from C. dubliniensis hybridized only to probe SEQ ID NO. 578, thereby demonstrating that each probe was species-specific.

EXAMPLE 9:

Specific identification of *Entamoeba histolytica*. Upon analysis of *tuf* (elongation factor 1 alpha) sequence data, it was possible to find four regions where

eukaryotic species have diverged. Primers TEntG38 (SEQ ID NO. 703), TEntG442 (SEQ ID NO. 704), TEntG534 (SEQ ID NO. 705), and TEntG768 (SEQ ID NO. 706) were designed so that SEQ ID NO. 703 could be paired with the three other primers. On PTC-200 thermocyclers (MJ Research), the cycling conditions for initial sensitivity and specificity testing were 3 min. at 94 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 55 °C, followed by terminal extension at 72 °C for 2 minutes. Detection of the PCR products was made by electrophoresis in agarose gels (2 %) containing 0.25 μg/ml of ethidium bromide. The three primer pairs could detect the equivalent of less than 200 *E. histolytica* genome copies. Specificity was tested using 0.5 ng of purified genomic DNA from a panel of microorganisms including *Babesia bovis, Babesia microtti, Candida albicans, Crithidia fasciculata, Leishmania major, Leishmania hertigi* and *Neospora caninum*. Only *E. histolytica* DNA could be amplified, thereby suggesting that the assay was species-specific.

EXAMPLE 10:

Sensitive identification of *Chlamydia trachomatis*. Upon analysis of *tuf* sequence data, it was possible to find two regions where *Chlamydia trachomatis* sequences remained conserved while other species have diverged. Primers Ctr82 (SEQ ID NO. 554) and Ctr249 (SEQ ID NO. 555) were designed. With the PTC-200 thermocyclers (MJ Research), the optimal cycling conditions for maximum sensitivity and specificity were determined to be 3 min. at 94 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 60 °C, followed by terminal extension at 72 °C for 2 minutes. Detection of the PCR products was made by electrophoresis in agarose gels (2 %) containing 0.25 μ g/ml of ethidium bromide. The assay could detect the equivalent of 8 *C. trachomatis* genome copies. Specificity was tested with 0.1 ng of purified genomic DNA from a panel of microorganisms including 22 species commonly encountered

in the vaginal flora (Bacillus subtilis, Bacteroides fragilis, Candida albicans, Clostridium difficile, Corynebacterium cervicis, Corynebacterium urealyticum, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Fusobacterium nucleatum, Gardnerella vaginalis, Haemophilus influenzae, Klebsiella oxytoca, Lactobacillus acidophilus, Peptococcus niger, Peptostreptococcus prevotii, Porphyromonas asaccharolytica, Prevotella melaninogenica, Propionibacterium acnes, Staphylococcus aureus, Streptococcus acidominimus, and Streptococcus agalactiae). Only C. trachomatis DNA could be amplified, thereby suggesting that the assay was species-specific.

EXAMPLE 11:

Genus-specific detection and identification of enterococci. Upon analysis of tuf sequence data and comparison with the repertory of tuf sequences, it was possible to find two regions where Enterococcus sequences remained conserved while other genera have diverged (Annex XVII). Primer pair Encg313dF and Encg599c (SEQ ID NOs. 1137 and 1136) was tested for its specificity by using purified genomic DNA from a panel of bacteria listed in Table 10. Using the PTC-200 thermocycler (MJ Research), the optimal cycling conditions for maximum sensitivity and specificity were determined to be 3 min. at 94 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 55 °C, followed by terminal extension at 72 °C for 2 minutes. Detection of the PCR products was made by electrophoresis in agarose gels (2 %) containing 0.25 μ g/ml of ethidium bromide. Visualization of the PCR products was made under UV at 254 nm. The 18 enterococcal species listed in Table 10 were all amplified efficiently. The only other species amplified were Abiotrophia adiacens, Gemella haemolysans and Gemella morbillorum, three gram-positive species. Sensitivity tested with several strains of E. casseliflavus, E. faecium, E. faecalis, E. flavescens and E. gallinarum and with one strain of each other Enterococcus species listed in Table 10 ranged from 1 to 10 copies of genomic DNA. The sequence variation

within the 308-bp amplicon was sufficient so that internal probes could be used to speciate the amplicon and differenciate enterococci from Abiotrophia adiacens, Gemella haemolysans and Gemella morbillorum, thereby allowing to achieve excellent specificity. Species-specific internal probes were generated for each of the clinically important species, E. faecalis (SEQ ID NO. 1174), E. faecium (SEQ ID NO. 602), and the group including E. casseliflavus, E. flavescens and E. gallinarum (SEQ ID NO. 1122) (Annex XVIII). The species-specific internal probes were able to differentiate their respective Enterococcus species from all other Enterococcus species. These assays are sensitive, specific and ubiquitous for those five Enterococcus species.

EXAMPLE 12:

Identification of the major bacterial platelets contaminants using tuf sequences with a multiplex PCR test. Blood platelets preparations need to be monitored for bacterial contaminations. The tuf sequences of 17 important bacterial contaminants of platelets were aligned. As shown in Annex XIX, analysis of these sequences allowed the design of PCR primers. Since in the case of contamination of platelet concentrates, detecting all species (not just the more frequently encountered ones) is desirable, perfect specificity of primers was not an issue in the design. However, sensitivity is important. That is why, to avoid having to put too much degeneracy, only the most frequent contaminants were included in primer design, knowing that the selected primers would anyway be able to amplify more species than the 17 used in the design because they target highly conserved regions of tuf sequences. Oligonucleotide sequences which are conserved in these 17 major bacterial contaminants of platelet concentrates were chosen (oligos Tplaq 769 and Tplaq 991, respectively SEQ ID NOs. 636 and 637) thereby permitting the detection of these bacterial species. However, sensitivity was slightly deficient with staphylococci. To ensure maximal sensitivity in the detection of all the more frequent bacterial contaminants, a multiplex assay also including oligonucleotide

primers targetting the *Staphylococcus* genera (oligos Stag 422, SEQ ID NO. 553; and Stag 765, SEQ ID NO. 575) was developed. The bacterial species detected with the assay are listed in Table 14.

The primer pairs, oligos SEQ ID NO. 636 and SEQ ID NO. 637 that give an amplification product of 245 pb, and oligos SEQ ID NO. 553 and SEQ ID NO. 575 that give an amplification product of 368 pb, were used simultaneously in the multiplex PCR assay. Detection of these PCR products was made on the LightCycler thermocycler (Idaho Technology) using SYBR® Green I (Molecular Probe Inc.). SYBR® Green I is a fluorescent dye that binds specifically to double-stranded DNA.

Fluorogenic detection of PCR products with the LightCycler was carried out using 1.0 μ M of both Tplag primers (SEQ ID NOs. 636-637) and 0.4 μ M of both TStaG primers (SEO ID NOs. 553 and 575), 2.5 mM MgCl₂, BSA 7.5 μ M, dNTP 0.2 mM (Pharmacia), 10 mM Tris-HCl (pH 8.3), 50 mM KCl, 0.5 U Taq DNA polymerase (Boerhinger Mannheim) coupled with TaqStartTM antibody (Clontech), and 0.07 ng of genomic DNA sample in a final volume of 7 μ l. The optimal cycling conditions for maximum sensitivity and specificity were 1 minute at 94 °C for initial denaturation, then forty-five cycles of three steps consisting of 0 second at 95 °C, 5 seconds at 60 °C and 9 seconds at 72 °C. Amplification was monitored during each elongation cycle by measuring the level of SYBR® Green I. However, real analysis takes place after PCR. Melting curves are done for each sample and transformation of the melting peak allows determination of Tm. Thus primer-dimer and specific PCR product are discriminated. With this assay, all prominent bacterial contaminants of platelet concentrates listed in Annex XIX and Table 14 were detected. Sensitivity tests were performed on the 9 most frequent bacterial contaminants of platelets. The detection limit was less than 20 genome copies for E. cloacae, B. cereus, S. choleraesuis and S. marcescens; less than 15 genome copies for P. aeruginosa; and 2 to 3 copies were detected for S. aureus, S.

epidermidis, E. coli and K. pneumoniae. Further refinements of assay conditions should increase sensitivity levels.

EXAMPLE 13:

The resolving power of the tuf and atpD sequences databases is comparable to the biochemical methods for bacterial identification. The present gold standard for bacterial identification is mainly based on key morphological traits and batteries of biochemical tests. Here we demonstrate that the use of tuf and atpD sequences combined with simple phylogenetic analysis of databases formed by these sequences is comparable to the gold standard. In the process of acquiring data for the tuf sequences, we sequenced the tuf gene of a strain that was given to us labelled as Staphylococcus hominis ATCC 35982. That tuf sequence (SEQ ID NO. 192) was incorporated into the tuf sequences database and subjected to a basic phylogenic analysis using the Pileup command from version 10 of the GCG package (Genetics Computer Group). This analysis indicated that SEQ ID NO. 192 is not associated with other S. hominis strains but rather with the S. warneri strains. The ATCC 35982 strain was sent to the reference laboratory of the Laboratoire de santé publique du Québec (LSPQ). They used the classic identification scheme for staphylococci (Kloos and Schleifer, 1975., J. Clin. Microbiol. 1:82-88). Their results shown that although the colonial morphology could correspond to S. hominis, the more precise biochemical assays did not. These assays included discriminant mannitol, mannose and ribose acidification tests as well as rapid and dense growth in deep thioglycolate agar. The LSPQ report identified strain ATCC 35982 as S. warneri which confirms our database analysis. The same thing happened for S. warneri (SEQ ID NO. 187) which had initially been identified as S. haemolyticus by a routine clinical laboratory using a low resolving power automated system (MicroScan, AutoScan-4TM). Again, the tuf and LSPQ analysis agreed on its identification as S. warneri. In numerous other instances, in the course of acquiring tuf and atpD sequence data from various species and genera,

analysis of our *tuf* and/or *atpD* sequence databases permitted the exact identification of mislabelled or erroneously identified strains. These results clearly demonstrate the usefulness and the high resolving power of our sequence-based identification assays using the *tuf* and *atpD* sequences databases.

EXAMPLE 14:

Detection of group B streptococci from clinical specimens.

Introduction

Streptococcus agalactiae, the group B streptococcus (GBS), is responsible for a severe illness affecting neonate infants. The bacterium is passed from the healthy carrier mother to the baby during delivery. To prevent this infection, it is recommended to treat expectant mothers susceptible of carrying GBS in their vaginal/anal flora. Carrier status is often a transient condition and rigorous monitoring requires cultures and classic bacterial identification weeks before delivery. To improve the detection and identification of GBS we developed a rapid, specific and sensitive PCR test fast enough to be performed right at delivery.

Materials and Methods

GBS clinical specimens. A total of 66 duplicate vaginal/anal swabs were collected from 41 consenting pregnant women admitted for delivery at the Centre Hospitalier Universitaire de Québec, Pavillon Saint-François d'Assise following the CDC recommendations. The samples were obtained either before or after rupture of membranes. The swab samples were tested at the Centre de Recherche en Infectiologie de l'Université Laval within 24 hours of collection. Upon receipt, one swab was cut and then the tip of the swab was added to GNS selective broth for identification of group B streptococci (GBS) by the standard culture methods

recommended by the CDC. The other swab was processed following the instruction of the IDI DNA extraction kit (Infectio Diagnotics (IDI) Inc.) prior to PCR amplification.

Oligonucleotides. PCR primers, Tsag340 (SEQ ID NO. 549) and Tsag552 (SEQ ID NO. 550) complementary to the regions of the *tuf* gene unique for GBS were designed based upon a multiple sequence alignment using our repertory of *tuf* sequences. Oligo primer analysis software (version 5.0) (National Biosciences) was used to analyse primers annealing temperature, secondary structure potential as well as mispriming and dimerization potential. The primers were synthesized using a model 391 DNA synthesizer (Applied Biosystems).

A pair of fluorescently labeled adjacent hybridization probes Sag465-F (SEQ ID NO. 583) and Sag436-C (SEQ ID NO. 582) were synthesized and purified by Operon Technologies. They were designed to meet the recommendations of the manufacturer (Idaho Technology) and based upon multiple sequence alignment analysis using our repertory of *tuf* sequences to be specific and ubiquitous for GBS. These adjacent probes, which are separated by one nucleotide, allow fluorescence resonance energy transfer (FRET), generating an increased fluorescence signal when both hybridized simultaneously to their target sequences. The probe SEQ ID NO. 583 was labeled with FITC in 3 prime while SEQ ID NO. 582 was labeled with Cy5 in 5 prime. The Cy5-labeled probes contained a 3'-blocking phosphate group to prevent extension of the probes during the PCR reactions.

PCR amplification. Conventional amplifications were performed either from 2 μl of a purified genomic DNA preparation or cell lysates of vaginal/anal specimens. The 20 μl PCR mixture contained 0.4 μM of each GBS-specific primer (SEQ ID NOs. 549-550), 200 μM of each deoxyribonucleotide (Pharmacia Biotech), 10 mM Tris-HCl (pH 9.0), 50 mM KCl, 0.1% Triton X-100, 2.5 mM MgCl₂, 3.3 mg/ml bovine serum albumin (BSA) (Sigma), and 0.5 U of *Taq* polymerase (Promega) combined with the TaqStartTM antibody (Clontech). The TaqStartTM antibody, which is a neutralizing monoclonal antibody of *Taq* DNA

polymerase, was added to all PCR reactions to enhance the efficiency of the amplification. The PCR mixtures were subjected to thermal cycling (3 min at 95 °C and then 40 cycles of 1 s at 95 °C, and 30 s at 62 °C with a 2-min final extension at 72 °C) with a PTC-200 DNA Engine thermocycler (MJ research). The PCR-amplified reaction mixture was resolved by agarose gel electrophoresis.

The LightCyclerTM PCR amplifications were performed with 1 μ l of a purified genomic DNA preparation or cell lysates of vaginal/anal specimens. The $10\mu l$ amplification mixture consisted of 0.4 μM each GBS-specific primer (SEQ ID NOs. 549-550), 200 μ M each dNTP, 0.2 μ M each fluorescently labeled probe (SEQ ID NOs. 582-583), 300 μ g/ml BSA (Sigma), and 1 μ l of 10x PC2 buffer (containing 50 mM Tris-HCl (pH 9.1), 16 mM ammonium sulfate, 3.5 mM Mg²⁺, and 150 μ g/ml BSA) and 0.5 U KlenTaq1TM (AB Peptides) coupled with TaqStartTM antibody (Clontech). KlenTaq1TM is a highly active and more heatstable DNA polymerase without 5'-exonuclease activity. This prevents hydrolysis of hybridized probes by the 5' to 3' exonuclease activity. A volume of 7 μ l of the PCR mixture was transferred into a composite capillary tube (Idaho Technology). The tubes were then centrifuged to move the reaction mixture to the tips of the capillaries and then cleaned with optical-grade methanol. Subsequently the capillaries were loaded into the carousel of a LC32 LightCyclerTM (Idaho Technology), an instrument that combines rapid-cycle PCR with fluorescence analysis for continuous monitoring during amplification. The PCR reaction mixtures were subjected to a denaturation step at 94 °C for 3 min followed by 45 cycles of 0 s at 94 °C, 20 s at 64 °C and 10 s at 72 °C with a temperature transition rate of 20 °C/s. Fluorescence signals were obtained at each cycle by sequentially positioning each capillary on the carousel at the focus of optical elements affiliated to the built-in fluorimeter for 100 milliseconds. Complete amplification and analysis required about 35 min.

Specificity and sensitivity tests. The specificity of the conventional and LightCyclerTM PCR assays was verified by using purified genomic DNA (0.1 ng/reaction) from a battery of ATCC reference strains representing 35 clinically

defectiva ATCC 49176, species (Abiotrophia gram-positive relevant Bisidobacterium breve ATCC 15700, Clostridium dissicile ATCC 9689, Corynebacterium urealyticum ATCC 43042, Enterococcus casseliflavus ATCC 25788, Enterococcus durans ATCC 19432, Enterococcus faecalis ATCC 29212, Enterococcus faecium ATCC 19434, Enterococcus gallinarum ATCC 49573, Enterococcus raffinosus ATCC 49427, Lactobacillus reuteri ATCC 23273, Lactococcus lactis ATCC 19435, Listeria monocytogenes ATCC 15313, Peptococcus niger ATCC 27731, Peptostreptococcus anaerobius ATCC 27337, Peptostreptococcus prevotii ATCC 9321, Staphylococcus aureus ATCC 25923, Staphylococcus epidermidis ATCC 14990, Staphylococcus haemolyticus ATCC 29970, Staphylococcus saprophyticus ATCC 15305, Streptococcus agalactiae ATCC 27591, Streptococcus anginosus ATCC 33397, Streptococcus bovis ATCC 33317, Streptococcus constellatus ATCC 27823, Streptococcus dysgalactiae ATCC 43078, Streptococcus gordonii ATCC 10558, Streptococcus mitis ATCC 33399, Streptococcus mutans ATCC 25175, Streptococcus oralis ATCC 35037, Streptococcus parauberis ATCC 6631, Streptococcus pneumoniae ATCC 6303, Streptococcus pyogenes ATCC 19615, Streptococcus salivarius ATCC 7073, Streptococcus sanguinis ATCC 10556, Streptococcus uberis ATCC 19436). These microbial species included 15 species of streptococci and many members of the normal vaginal and anal floras. In addition, 40 GBS isolates of human origin, whose identification was confirmed by a latex agglutination test (Streptex, Murex), were also used to evaluate the ubiquity of the assay.

For determination of the sensitivities (i.e., the minimal number of genome copies that could be detected) for conventional and LightCyclerTM PCR assays, serial 10-fold or 2-fold dilutions of purified genomic DNA from 5 GBS ATCC strains were used.

Results

Evaluation of the GBS-specific conventional and LightCyclerTM PCR assays. The specificity of the two assays demonstrated that only DNAs from GBS

strains could be amplified. Both PCR assays did not amplify DNAs from any other bacterial species tested including 14 streptococcal species other than GBS as well as phylogenetically related species belonging to the genera *Enterococcus*, *Peptostreptococcus* and *Lactococcus*. Important members of the vaginal or anal flora, including coagulase-negative staphylococci, *Lactobacillus* sp., and *Bacteriodes* sp. were also negative with the GBS-specific PCR assay. The LightCyclerTM PCR assays detected only GBS DNA by producing an increased fluorescence signal which was interpreted as a positive PCR result. Both PCR methods were able to amplify all of 40 GBS clinical isolates, showing a perfect correlation with the phenotypic identification methods.

The sensitivity of the assay was determined by using purified genomic DNA from the 5 ATCC strains of GBS. The detection limit for all of these 5 strains was one genome copy of GBS. The detection limit of the assay with the LightCyclerTM was 3.5 fg of genomic DNA (corresponding to 1-2 genome copies of GBS). These results confirmed the high sensitivity of our GBS-specific PCR assay.

Direct Detection of GBS from vaginal/anal specimens. Among 66 vaginal/anal specimens tested, 11 were positive for GBS by both culture and PCR. There was one sample positive by culture only. The sensitivity of both PCR methods with vaginal/anal specimens for identifying colonization status in pregnant women at delivery was 91.7% when compared to culture results. The specificity and positive predictive values were both 100% and the negative predictive value was 97.8%. The time for obtaining results was approximately 45 min for LightCyclerTM PCR, approximately 100 min for conventional PCR and 48 hours for culture.

Conclusion

We have developed two PCR assays (conventional and LightCyclerTM) for the detection of GBS, which are specific (i.e., no amplification of DNA from a variety of bacterial species other than GBS) and sensitive (i.e., able to detect around 1

genome copy for several reference ATCC strains of GBS). Both PCR assays are able to detect GBS directly from vaginal/anal specimens in a very short turnaround time. Using the real-time PCR assay on LightCyclerTM, we can detect GBS carriage in pregnant women at delivery within 45 minutes.

EXAMPLE 15:

Simultaneous detection and identification of Streptococcus pyogenes and its pyrogenic exotoxin A. The rapid detection of Streptococcus pyogenes and of its pyrogenic exotoxin A is of clinical importance. We developed a multiplex assay which permits the detection of strains of S. pyogenes carrying the pyrogenic toxin A gene, which is associated with scarlet fever and other pathologies. In order to specifically detect S. pyogenes, nucleotide sequences of the pyrrolidone carboxylyl peptidase (pcp) gene were aligned to design PCR primers Spy291 (SEQ ID NO. 1211) and Spy473 (SEQ ID NO. 1210). Next, we designed primers for the specific detection of the pyrogenic exotoxin A. Nucleotide sequences of the speA gene, carried on the bacteriophage T12, were aligned as shown in Annex XXIII to design PCR primers Spytx814 (SEQ ID NO. 994) and Spytx 927 (SEQ ID NO. 995).

The primer pairs: oligos SEQ ID NOs. 1210-1211, yielding an amplification product of 207 bp, and oligos SEQ ID NOs. 994-995, yielding an amplification product of 135 bp, were used in a multiplex PCR assay.

PCR amplification was carried out using 0.4 μ M of both pairs of primers, 2.5 mM MgCl₂, BSA 0.05 μ M, dNTP 0.2 μ M (Pharmacia), 10mM Tris-HCl (pH 9.0), 0.1% Triton X-100, 2.5 mM MgCl₂, 0.5 U *Taq* DNA polymerase (Promega) coupled with TaqStartTM antibody (Clontech Laboratories Inc.), and 1 μ l of genomic DNA sample in a final volume of 20 μ l. PCR amplification was performed using a PTC-200 thermal cycler (MJ Research). The optimal cycling conditions for maximum specificity and sensitivity were 3 minutes at 94 °C for

initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 63 °C, followed by a final step of 2 minutes at 72 °C. Detection of the PCR products was made by electrophoresis in agarose gels (2 %) containing 0.25 μ g/ml of ethidium bromide. Visualization of the PCR products was made under UV at 254 nm.

The detection limit was less than 5 genome copies for both *S. pyogenes* and its pyrogenic exotoxin A. The assay was specific for pyrogenic exotoxin A-producing *S. pyogenes*: strains of the 27 other species of *Streptococcus* tested, as well as 20 strains of various gram-positive and gram-negative bacterial species were all negative.

A similar approach was used to design an alternative set of *speA*-specific primers (SEQ ID NOs. 996 to 998, see Annex XXIV). In addition, another set of primers based on the *tuf* gene (SEQ ID NOs. 999 to 1001, see Annex XXV) could be used to specifically detect *Streptococcus pyogenes*.

EXAMPLE 16:

Real-time detection and identification of Shiga toxin-producing bacteria. Shiga toxin-producing Escherichia coli and Shigella dysenteriae cause bloody diarrhea. Currently, identification relies mainly on the phenotypic identification of S. dysenteriae and E. coli serotype O157:H7. However, other serotypes of E. coli are increasingly found to be producers of type 1 and/or type 2 Shiga toxins. Two pairs of PCR primers targeting highly conserved regions present in each of the Shiga toxin genes stx_1 and stx_2 were designed to amplify all variants of those genes (see Annexes XXVI and XXVII). The first primer pair, oligonucleotides 1SLT224 (SEQ ID NO. 1081) and 1SLT385 (SEQ ID NO. 1080), yields an amplification product of 186 bp from the stx_1 gene. For this amplicon, the 1SLTB1-Fam (SEQ ID NO. 1084) molecular beacon was designed for the specific detection of stx_1

using the fluorescent label 6-carboxy-fluorescein. The 1SltS1-FAM (SEQ ID NO. 2012) molecular scorpion was also designed as an alternate way for the specific detection of stx_1 . A second pair of PCR primers, oligonucleotides 2SLT537 (SEQ ID NO. 1078) and 2SLT678b (SEQ ID NO. 1079), yields an amplification product of 160 bp from the stx_2 gene. Molecular beacon 2SLTB1-Tet (SEQ ID NO. 1085) was designed for the specific detection of stx_2 using the fluorescent label 5-tetrachloro-fluorescein. Both primer pairs were combined in a multiplex PCR assay.

PCR amplification was carried out using 0.8 μM of primer pair SEQ ID NOs. 1080-1081, 0.5 μM of primer pair SEQ ID NOs. 1078-1079, 0.3 μM of each molecular beacon, 8 mM MgCl₂, 490 μg/mL BSA, 0.2 mM dNTPs (Pharmacia), 50 mM Tris-HCl, 16 mM NH₄SO₄, 1X TaqMaster (Eppendorf), 2.5 U KlenTaq1 DNA polymerase (AB Peptides) coupled with TaqStartTM antibody (Clontech Laboratories Inc.), and 1 μl of genomic DNA sample in a final volume of 25 μl. PCR amplification was performed using a SmartCycler thermal cycler (Cepheid). The optimal cycling conditions for maximum sensitivity and specificity were 60 seconds at 95 °C for initial denaturation, then 45 cycles of three steps consisting of 10 seconds at 95 °C, 15 seconds at 56 °C and 5 seconds at 72 °C. Detection of the PCR products was made in real-time by measuring the fluorescent signal emitted by the molecular beacon when it hybridizes to its target at the end of the annealing step at 56 °C.

The detection limit was the equivalent of less than 5 genome copies. The assay was specific for the detection of both toxins, as demonstrated by the perfect correlation between PCR results and the phenotypic characterization performed using antibodies specific for each Shiga toxin type. The assay was successfully performed on several Shiga toxin-producing strains isolated from various geographic areas of the world, including 10 O157:H7 E. coli, 5 non-O157:H7 E. coli and 4 S. dysenteriae.

EXAMPLE 17:

Development of a PCR assay for the detection and identification of staphylococci at genus and species levels and its associated mecA gene. The Staphylococcusspecific PCR primers described in Example 7 (SEQ ID NOs. 553 and 575) were used in multiplex with the mecA-specific PCR primers and the S. aureus-specific primers described in our assigned US patent no. 5,994,066 (SEQ ID NOs. 261 and 262 for mecA and SEQ ID NOs. 152 and 153 for S.aureus in the said patent). Sequence alignment analysis of 10 publicly available mecA gene sequences allowed to design an internal probe specific to mecA (SEQ ID NO. 1177). An internal probe was also designed for the S. aureus-specific amplicon (SEQ ID NO 1234). PCR amplification and agarose gel electrophoresis of the amplified products were performed as described in Example 7, with the exception that 0.4 μM (each) of the two Staphylococcus-specific primers (SEQ ID NOs. 553 and 575) and 0.4 μM (each) of the mecA-specific primers and 0.4 μM (each) of the S. aureusspecific primers were used in the PCR mixture. The specificity of the multiplex assay with 40-cycle PCR protocols was verified by using purified genomic DNA from five methicillin-resistant and fifteen methicillin-sensitive staphylococcal strains. The sensitivity of the multiplex assay with 40-cycle PCR protocols was determined by using purified genomic DNA from twenty-three-methicillinresistant and twenty-eight methicillin-sensitive staphylococcal strains. The detection limit was 2 to 10 genome copies of genomic DNA, depending on the staphylococcal species tested. Furthermore, the mecA-specific internal probe, the S. aureus-specific internal probe and the coagulase-negative staphylococci-specific internal probe (described in Example 7) were able to recognize twenty-three methicillin-resistant staphylococcal strains and twenty-eight methicillin-sensitive staphylococcal strains with high sensitivity and specificity.

The format of the assay is not limited to the one described above. A person skilled in the art could adapt the assay for different formats such as PCR with real-time detection using molecular beacon probes. Molecular beacon probes designed to be used in this assay include, but are not limited to, SEQ ID NO. 1232 for detection of the S. aureus-specific amplicon, SEQ ID NO. 1233 for detection of coagulase-negative staphylococci and SEQ ID NO. 1231 for detection of mecA.

Alternatively, a multiplex PCR assay containing the Staphylococcus-specific PCR primers described in Example 7 (SEQ ID NOs. 553 and 575) and the mecAspecific PCR primers described in our assigned US patent no. 5,994,066 (SEQ ID NOs. 261 and 262 in the said patent) were developed. PCR amplification and agarose gel electrophoresis of the amplified products were performed as described in Example 7, with the exception that 0.4 μ M (each) of the Staphylococcus-specific primers (SEQ ID NOs. 553 and 575) and 0.4 μM (each) of the mecA-specific primers described in our assigned US patent no. 5,994,066 (SEQ ID NOs. 261 and 262 in the said patent) were used in the PCR mixture. The sensitivity of the multiplex assay with 40-cycle PCR protocols was determined by using purified genomic DNA from two methicillin-resistant and five methicillin-sensitive staphylococcal strains. The detection limit was 2 to 5 copies of genomic DNA, depending on the staphylococcal species tested. The specificity of the multiplex PCR assay coupled with capture-probe hybridization was tested with two strains of methicillin-resistant S. aureus, two strains of methicillin-sensitive S. aureus and seven strains of methicillin-sensitive coagulase-negative staphylococci. The mecAspecific internal probe (SEQ ID NO. 1177) and the S. aureus-specific internal probe (SEQ ID NO. 587) described in Example 7 were able to recognize all the strains with high specificity showing a perfect correlation with susceptibility to methicillin. The sensitivity of the PCR assay coupled with capture-probe hybridization was tested with one strain of methicillin-resistant S. aureus. The detection limit was around 10 copies of genomic DNA.

EXAMPLE 18:

Sequencing of pbp1a, pbp2b and pbp2x genes of Streptoccoccus pneumoniae. Penicillin resistance in Streptococcus pneumoniae involves the sequential alteration of up to five penicillin-binding proteins (PBPs) 1A, 1B, 2A, 2X and 2B in such a way that their affinity is greatly reduce toward the antibiotic molecule. The altered PBP genes have arisen as the result of interspecies recombination events from related streptococcal species. Among the PBPs usually found in S. pneumoniae, PBPs 1A, 2B, and 2X play the most important role in the development of penicillin resistance. Alterations in PBP 2B and 2X mediate low-level resistance to penicillin while additional alterations in PBP 1A plays a significant role in full penicillin resistance.

In order to generate a database for pbp sequences that can be used for design of primers and/or probes for the specific and ubiquitous detection of β-lactam resistance in S. pneumoniae, pbp1a, pbp2b and pbp2x DNA fragments sequenced by us or selected from public databases (GenBank and EMBL) from a variety of S. pneumoniae strains were used to design oligonucleotide primers. This database is essential for the design of specific and ubiquitous primers and/or probes for detection of \beta-lactam resistance in S. pneumoniae since the altered PBP 1A, PBP 2B and PBP 2X of β-lactam resistant S. pneumoniae are encoded by mosaic genes with numerous sequence variations among resistant isolates. The PCR primers were located in conserved regions of pbp genes and were able to amplify pbpla, pbp2b, and pbp2x sequences of several strains of S. pneumoniae having various levels of resistance to penicillin and third-generation cephalosporins. Using primer pairs SEQ ID NOs. 1125 and 1126, SEQ ID NOs. 1142 and 1143, SEQ ID NOs. 1146 and 1147, it was possible to amplify and determine pbpla sequences SEQ ID NOs. 1004-1018, 1648, 2056-2060 and 2062-2064, pbp2b sequences SEQ ID NOs. 1019-1033, and pbp2x sequences SEQ ID NOs. 1034-1048. Six other PCR primers

(SEQ ID NOs. 1127-1128, 1144-1145, 1148-1149) were also designed and used to complete the sequencing of pbp1a, pbp2b and pbp2x amplification products. The described primers (SEQ ID NOs. 1125 and 1126, SEQ ID NOs. 1142 and 1143, SEQ ID NOs. 1146 and 1147, SEQ ID NOs. 1127-1128, 1144-1145, 1148-1149) represent a powerful tool for generating new pbp sequences for design of primers and/or probes for detection of β -lactam resistance in S. pneumoniae.

EXAMPLE 19:

Sequencing of hexA genes of Streptococcus species. The hexA sequence of S. pneumoniae described in our assigned US patent no. 5,994,066 (SEQ ID NO. 31 in the said patent, SEQ ID NO. 1183 in the present application) allowed the design of a PCR primer (SEQ ID NO. 1182) which was used with primer Spn1401 described in our assigned US patent no. 5,994,066 (SEQ ID NO. 156 in the said patent, SEQ ID NO. 1179 in the present application) to generate a database for hexA sequences that can be used to design primers and/or probes for the specific identification and detection of S. pneumoniae (Annex XLII). Using primers SEQ ID NO. 1179 and SEQ ID NO. 1182 (Annex XLII), it was possible to amplify and determine the hexA sequence from S. pneumoniae (4 strains) (SEQ ID NOs. 1184-1187), S. mitis (three strains) (SEQ ID NOs. 1189-1191) and S. oralis (SEQ ID NO. 1188).

EXAMPLE 20:

Development of multiplex PCR assays coupled with capture probe hybridization for the detection and identification of *Streptococcus pneumoniae* and its penicillin resistance genes.

Two different assays were developed to identify S. pneumoniae and its susceptibility to penicillin.

ASSAY I:

Bacterial strains. The specificity of the multiplex PCR assay was verified by using a panel of ATCC (American Type Culture Collection) reference strains consisting of 33 gram-negative and 67 gram-positive bacterial species (Table 13). In addition, a total of 98 strains of *S. pneumoniae*, 16 strains of *S. mitis* and 3 strains of *S. oralis* from the American Type Culture Collection, the microbiology laboratory of the Centre Hospitalier Universitaire de Québec, Pavillon Centre Hospitalier de l'Université Laval (CHUL), (Ste-Foy, Québec, Canada), the Laboratoire de santé publique du Québec, (Sainte-Anne-de-Bellevue, Québec, Canada), the Sunnybrook and Women's College Health Sciences Centre (Toronto, Canada), the Infectious Diseases Section, Department of Veterans Affairs Medical Center, (Houston, USA) were also tested to further validate the *Streptococcus pneumoniae*-specific PCR assay. The penicillin MICs (minimal inhibitory concentrations) were measured by the broth dilution method according to the recommended protocol of NCCLS.

PCR primers and internal probes. The analysis of hexA sequences from a variety of streptococcal species from the publicly avalaible hexA sequence and from the database described in Example 19 (SEQ ID NOs. 1184-1191) allowed the selection of a PCR primer specific to S. pneumoniae, SEQ ID NO. 1181. This primer was used with the S. pneumoniae-specific primer SEQ ID NO. 1179 to generate an amplification product of 241 bp (Annex XLII). The PCR primer SEQ ID NO. 1181 is located 127 nucleotides downstream on the hexA sequence compared to the original S. pneumoniae-specific PCR primer Spn1515 described in our assigned US patent no. 5,994,066 (SEQ ID NO. 157 in the said patent). These modifications were done to ensure the design of the S. pneumoniae-specific internal probe according to the new hexA sequences of several streptococcal species from the database described in Example 19 (SEQ ID NOs. 1184-1191).

The analysis of pbp1a sequences from *S. pneumoniae* strains with various levels of penicillin resistance from public databases and from the database described in Example 18 allowed the identification of amino acid substitutions Ile-459 to Met and Ser-462 to Ala that occur in isolates with high-level penicillin resistance (MICs $\geq 1 \mu g/ml$), and amino acid substitutions Ser-575 to Thr, Gln-576 to Gly and Phe-577 to Tyr that are common to all penicillin-resistant isolates with MICs \geq 0.25 $\mu g/ml$. As shown in Annex XXXI, PCR primer pair SEQ ID NOs. 1130 and 1131 were designed to detect high-level penicillin resistance (MICs $\geq 1 \mu g/ml$), whereas PCR primer pair SEQ ID NOs. 1129 and 1131 were designed to detect intermediate- and high-level penicillin resistance (MICs \geq 0.25 $\mu g/ml$).

The analysis of hexA sequences from the publicly avalaible hexA sequence and from the database described in Example 19 allowed the design of an internal probe specific to S. pneumoniae (SEQ ID NO. 1180) (Annex XLII). The range of mismatches between the S. pneumoniae-specific 241-bp amplicon was from 2 to 5, in the middle of the 19-bp probe. The analysis of pbpla sequences from public databases and from the database described in Example 18 allowed the design of five internal probes containing all possible mutations to detect the high-level penicillin resistance 383-bp amplicon (SEQ ID NOs. 1197, 1217-1220). Alternatively, two other internal probes (SEQ ID NOs. 2024-2025) can also be used to detect the high-level penicillin resistance 383-bp amplicon. Five internal probes containing all possible mutations to detect the 157-bp amplicon which includes intermediate- and high-level penicillin resistance were also designed (SEQ ID NOs. 1094, 1192-1193, 1214 and 1216). Design and synthesis of primers and probes, and detection of the probe hybridization were performed as described in Example 7. Annex XXXI illustrates one of the internal probe for detection of the high-level penicillin resistance 383-bp amplicon (SEQ ID NO. 1197) and one of the internal probe for detection of the intermediate- and high-level penicillin resistance 157-bp amplicon (SEQ ID NO. 1193).

PCR amplification. For all bacterial species, amplification was performed from purified genomic DNA using a PTC-200 thermocycler (MJ Research). 1 μ l of genomic DNA at 0.1 ng/ μ l, or 1 μ l of a bacterial lysate, was transferred to a 19 μ l PCR mixture. Each PCR reaction contained 50 mM KCl, 10 mM Tris-HCl (H 9.0), 0.1% Triton X-100, 2.5 mM MgCl₂, 0.1 μ M (each) of the *S. pneumoniae*-specific primers SEQ ID NO. 1179 and SEQ ID NO. 1181, 0.2 μ M of primer SEQ ID NO. 1129, 0.7 μ M of primer SEQ ID NO. 1131, and 0.6 μ M of primer SEQ ID NO. 1130, 0.05 mM bovine serum albumin (BSA), and 0.5 U *Taq* polymerase (Promega) coupled with TaqStartTM antibody. In order to generate Digoxigenin (DIG)-labeled amplicons for capture probe hybridization, 0.1X PCR DIG labeling four deoxynucleoside triphosphates mix (Boehringer Mannheim GmbH) was used for amplification.

For determination of the sensitivity of the PCR assays, 10-fold dilutions of purified genomic DNA were used to determine the minimal number of genome copies which can be detected.

Capture probe hybridization. The DIG-labeled amplicons were hybridized to the capture probes bound to 96-well plates. The plates were incubated with anti-DIG-alkaline phosphatase and the chemiluminescence was measured by using a luminometer (MLX, Dynex Technologies Inc.) after incubation with CSPD and recorded as Relative Light Unit (RLU). The RLU ratio of tested sample with and without captures probes was then calculated. A ratio \geq 2.0 was defined as a positive hybridization signal. All reactions were performed in duplicate.

Results

Amplifications with the multiplex PCR assay. The specificity of the assay was assessed by performing 40-cycle PCR amplifications with the panel of grampositive (67 species from 12 genera) and gram-negative (33 species from 17

genera) bacterial species listed in Table 13. All bacterial species tested other than S. pneumoniae were negative except S. mitis and S. oralis. Ubiquity tests were performed using a collection of 98 S. pneumoniae strains including high-level penicillin resistance (n=53), intermediate resistance (n=12) and sensitive (n=33) strains. There was a perfect correlation between PCR and standard susceptibility testing for 33 penicillin-sensitive isolates. Among 12 S. pneumoniae isolates with intermediate penicillin resistance based on susceptibility testing, 11 had intermediate resistance based on PCR, but one S. pneumoniae isolate with penicillin MIC of 0.25 μ g/ml showed a high-level penicillin resistance based on susceptibility testing, 51 had high-level penicillin resistance based on PCR but two isolates with penicillin MIC > 1 μ g/ml showed an intermediate penicillin resistance based on genotyping. In general, there was a good correlation between the genotype and classical culture method for bacterial identification and susceptibility testing.

The sensitivity of the S. pneumoniae-specific assay with 40-cycle PCR protocols was determined by using purified genomic DNA from 9 isolates of S. pneumoniae. The detection limit was around 10 copies of genomic DNA for all of them.

Post-PCR hybridization with internal probes. The specificity of the multiplex PCR assay coupled with capture-probe hybridization was tested with 98 strains of S. pneumoniae, 16 strains of S. mitis and 3 strains of S. oralis. The internal probe specific to S. pneumoniae (SEQ ID NO. 1180) detected all 98 S. pneumoniae strains but did not hybridize to the S. mitis and S. oralis amplicons. The five internal probes specific to the high-level resistance amplicon (SEQ ID NOs. 1197, 1217-1220) detected all amplification patterns corresponding to high-level resistance. The two S. pneumoniae strains with penicillin MIC > 1 μ g/ml that showed an intermediate penicillin resistance based on PCR amplification were also intermediate resistance based on probe hybridization. Similarly, among 12 strains

with intermediate-penicillin resistance based on susceptibility testing, 11 showed intermediate-penicillin resistance based on hybridization with the five internal probes specific to the intermediate and high-level resistance amplicon (SEQ ID NOs. 1094, 1192-1193, 1214 and 1216). The strain described above having a penicillin MIC of 0.25 µg/ml which was high-level penicillin resistance based on PCR amplification was also high-level resistance based on probe hybridization. In summary, the combination of the multiplex PCR and hybridization assays results in a highly specific test for the detection of penicillin-resistant *Streptococcus pneumoniae*.

ASSAY II:

Bacterial strains. The specificity of the multiplex PCR assay was verified by using the same strains as those used for the development of Assay I. The penicillin MICs (minimal inhibitory concentrations) were measured by the broth dilution method according to the recommended protocol of NCCLS.

PCR primers and internal probes. The analysis of pbp1a sequences from S. pneumoniae strains with various levels of penicillin resistance from public databases and from the database described in Example 18 allowed the design of two primers located in the constant region of pbp1a. PCR primer pair (SEQ ID NOs. 2015 and 2016) was designed to amplify a 888-bp variable region of pbp1a from all S. pneumoniae strains. A series of internal probes were designed for identification of the pbp1a mutations associated with penicillin resistance in S. pneumoniae. For detection of high-level penicillin resistance (MICs $\geq 1 \mu g/ml$), three internal probes were designed (SEQ ID NOs. 2017-2019). Alternaltively, ten other internal probes were designed that can also be used for detection of high-level resistance within the 888-bp pbp1a amplicon: (1) three internal probes for identification of the amino acid substitutions Thr-371 to Ser or Ala within the motif S370TMK (SEQ ID NOs. 2031-2033); (2) two internal probes for detection

of the amino acid substitutions Ile-459 to Met and Ser-462 to Ala near the motif S428RN (SEQ ID NOs. 1135 and 2026); (3) two internal probes for identification of the amino acid substitutions Asn-443 to Asp (SEQ ID NOs. 1134 and 2027); and (4) three internal probes for detection of all sequence variations within another region (SEQ ID NOs. 2028-2030). For detection of high-level and intermediate penicillin resistance (MICs \geq 0.25 µg/ml), four internal probes were designed (SEQ ID NOs. 2020-2023). Alternatively, six other internal probes were designed for detection of the four consecutive amino acid substitutions T574SQF to A574TGY near the motif K557TG (SEQ ID NOs. 2034-2039) that can also be used for detection of intermediate- and high-level resistance within the 888-bp pbp1a amplicon.

PCR amplification. For all bacterial species, amplification was performed from purified genomic DNA using a PTC-200 thermocycler (MJ Research). 1 μ l of genomic DNA at 0.1 ng/ μ l, or 1 μ l of a bacterial lysate, was transferred to a 19 μ l PCR mixture. Each PCR reaction contained 50 mM KCl, 10 mM Tris-HCl (pH 9.0), 0.1% Triton X-100, 2.5 mM MgCl₂, 0.08 μ M (each) of the *S. pneumoniae*-specific primers SEQ ID NO. 1179 and SEQ ID NO. 1181, 0.4 μ M of the *pbp1a*-specific primer SEQ ID NO. 2015, 1.2 μ M of *pbp1a*-specific primer SEQ ID NO. 2016, 0.05 mM bovine serum albumin (BSA), and 0.5 U *Taq* polymerase (Promega) coupled with TaqStartTM antibody. In order to generate Digoxigenin (DIG)-labeled amplicons for capture probe hybridization, 0.1X PCR DIG labeling four deoxynucleoside triphosphates mix (Boehringer Mannheim GmbH) was used for amplification.

For determination of the sensitivities of the PCR assays, 10-fold dilutions of purified genomic DNA were used to determine the minimal number of genome copies which can be detected.

Capture probe hybridization. The DIG-labeled amplicons were hybridized to the capture probes bound to 96-well plates as described for Assay I.

Results

Amplifications with the multiplex PCR assay. The specificity of the assay was assessed by performing 40-cycle PCR amplifications with the panel of grampositive (67 species from 12 genera) and gram-negative (33 species from 17 genera) bacterial species listed in Table 13. All bacterial species tested other than S. pneumoniae were negative except S. mitis and S. oralis. Ubiquity tests were performed using a collection of 98 S. pneumoniae strains including high-level penicillin resistance (n=53), intermediate resistance (n=12) and sensitive (n=33) strains. All the above S. pneumoniae strains produced the 888-bp amplicon corresponding to pbp1a and the 241-bp fragment corresponding to hexA.

The sensitivity of the *S. pneumoniae*-specific assay with 40-cycle PCR protocols was determined by using purified genomic DNA from 9 isolates of *S. pneumoniae*. The detection limit was around 10 copies of genomic DNA for all of them.

Post-PCR hybridization with internal probes. The specificity of the multiplex PCR assay coupled with capture-probe hybridization was tested with 98 strains of *S. pneumoniae*, 16 strains of *S. mitis* and 3 strains of *S. oralis*. The internal probe specific to *S. pneumoniae* (SEQ ID NO. 1180) detected all 98 *S. pneunoniae* strains but did not hybridize to the *S. mitis* and *S. oralis* amplicons. The three internal probes (SEQ ID NOs 2017-2019) specific to high-level resistance detected all the 43 strains with high-level penicillin resistance based on susceptibility testing. Among 12 isolates with intermediate-penicillin resistance based on hybridization with 4 internal probes (SEQ ID NOs. 2020-2023) and one strain

having penicillin MIC of $0.25 \mu g/ml$ was misclassified as high-level penicillin resistance. In summary, the combination of the multiplex PCR and hybridization assays results in a highly specific test for the detection of penicillin-resistant Streptococcus pneumoniae.

EXAMPLE 21:

Sequencing of the vancomycin resistance vanA, vanC1, vanC2 and vanC3 genes. The publicly available sequences of the vanH-vanA-vanX-vanY locus of transposon Tn1546 from E. faecalis, vanC1 sequence from one strain of E. gallinarum, vanC2 and vanC3 sequences from a variety of E. casseliflavus and E. flavescens strains, respectively, allowed the design of PCR primers able to amplify the vanA, vanC1, vanC2 and vanC3 sequences of several Enterococcus species. Using primer pairs van6877 and van9106 (SEQ ID NOs. 1150 and 1155), vanC1-122 and vanC1-1315 (SEQ ID NOs. 1110 and 1109), and vanC2C3-1 and vanC2C3-1064 (SEQ ID NOs. 1108 and 1107), it was possible to amplify and determine vanA sequences SEQ ID NOs. 1049-1057, vanC1 sequences SEQ ID NOs. 1058-1059, vanC2 sequences SEQ ID NOs. 1060-1063 and vanC3 sequences SEQ ID NOs. 1151-1154) were also designed and used to complete the sequencing of vanA amplification products.

EXAMPLE 22:

Development of a PCR assay for the detection and identification of enterococci at genus and species levels and its associated resistance genes vanA and vanB. The comparison of vanA and vanB sequences revealed conserved regions allowing the design of PCR primers specific to both vanA and vanB sequences (Annex XXXVIII). The PCR primer pair vanAB459 and vanAB830R (SEQ ID NOs. 1112 and 1111) was used in multiplex with the Enterococcus-specific primers Encg313dF and Encg599c (SEQ ID NOs. 1137 and 1136) described in Example

11. Sequence alignment analysis of vanA and vanB sequences revealed regions suitable for the design of internal probes specific to vanA (SEQ ID NO. 1170) and vanB (SEQ ID NO. 1171). PCR amplification and agarose gel electropheresis of the amplified products were performed as described in Example 11. The optimal cycling conditions for maximum sensitivity and specificity were found to be 3 min. at 94 °C, followed by forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 62°C, plus a terminal extension at 72 °C for 2 minutes. The specificity of the multiplex assay with 40-cycle PCR was verified by using 0.1 nanogram of purified genomic DNA from a panel of bacteria listed in Table 10. The sensitivity of the multiplex assay with 40-cycle PCR was verified with three strains of E. casseliflavus, eight strains of E. gallinarum, two strains of E. flavescens, two vancomycin-resistant strains of E. faecalis and one vancomycinsensitive strain of E. faecalis, three vancomycin-resistant strains of E. faecium, one vancomycin-sensitive strain of E. faecium and one strain of each of the other enterococcal species listed in Table 10. The detection limit was 1 to 10 copies of genomic DNA, depending on the enterococcal species tested. The vanA- and vanBspecific internal probes (SEQ ID NOs. 1170 and 1171), as well as the E. faecalisand E. faecium-specific internal probes (SEQ ID NOs. 1174 and 602) and the internal probe specific to the group including E. casseliflavus, E. gallinarum and E. flavescens (SEQ ID NO. 1122) described in Example 11, were able to recognize vancomycin-resistant enterococcal species with high sensitivity, specificity and ubiquity showing a perfect correlation between the genotypic and phenotypic analysis.

The format of the assay is not limited to the one described above. A person skilled in the art could adapt the assay for different formats such as PCR with real-time detection using molecular beacon probes. Molecular beacon probes designed to be used in this assay include, but are not limited to, SEQ ID NO. 1236 for the detection of *E. faecalis*, SEQ ID NO. 1235 for the detection of *E. faecium*, SEQ ID NO. 1240 for the detection of *vanA*, and SEQ ID NO. 1241 for the detection of *vanB*.

EXAMPLE 23:

Development of a multiplex PCR assay for detection and identification of vancomycin-resistant Enterococcus faecalis, Enterococcus faecium and the group including Enterococcus gallinarum, Enterococcus casseliflavus, and Enterococcus flavescens. The analysis of vanA and vanB sequences revealed conserved regions allowing design of a PCR primer pair (SEQ ID NOs. 1089 and 1090) specific to vanA sequences (Annex XXVIII) and a PCR primer pair (SEQ ID NOs. 1095 and 1096) specific to vanB sequences (Annex XXIX). The vanA-specific PCR primer pair (SEO ID NOs. 1089 and 1090) was used in multiplex with the vanB-specific PCR primer pair described in our assigned US patent 5,994,066 (SEQ ID NOs. 1095 and 1096 in the present patent and SEQ ID NOs. 231 and 232 in the said patent). The comparison of vanC1, vanC2 and vanC3 sequences revealed conserved regions allowing design of PCR primers (SEQ ID NOs. 1101 and 1102) able to generate a 158-bp amplicon specific to the group including E. gallinarum, E. casseliflavus and E. flavescens (Annex XXX). The vanC-specific PCR primer pair (SEQ ID NOs. 1101 and 1102) was used in multiplex with the E. faecalisspecific PCR primer pair described in our assigned US patent 5,994,066 (SEQ ID NOs. 40 and 41 in the said patent) and with the E. faecium-specific PCR primer pair described in our patent publication WO98/20157 (SEQ ID NOs. 1 and 2 in the said publication). For both multiplexes, the optimal cycling conditions for maximum sensitivity and specificity were found to be 3 min. at 94 °C, followed by forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 58 °C, plus a terminal extension at 72 °C for 2 minutes. Detection of the PCR products was made by electrophoresis in agarose gels (2 %) containing 0.25 μ g/ml of ethidium bromide. The vanA-specific PCR primer pair (SEQ ID NOs. 1089 and 1090), the vanB-specific primer pair (SEQ ID NOs. 1095 and 1096) and the vanCspecific primer pair (SEQ ID NOs. 1101 and 1102) were tested for their specificity by using 0.1 nanogram of purified genomic DNA from a panel of 5 vancomycin-

sensitive Enterococcus species, 3 vancomycin-resistant Enterococcus species, 13 other gram-positive bacteria and one gram-negative bacterium. Specificity tests were performed with the E. faecium-specific PCR primer pair described in our patent publication WO98/20157 (SEQ ID NOs. 1 and 2 in the said publication) and with the E. faecalis-specific PCR primer pair described in our assigned US patent 5,994,066 (SEQ ID NOs. 40 and 41 in the said patent) on a panel of 37 gram-positive bacterial species. All Enterococcus strains were amplified with high specificity showing a perfect correlation between the genotypic and phenotypic analysis. The sensitivity of the assays was determined for several strains of E. gallinarum, E. casseliflavus, E. flavescens and vancomycin-resistant E. faecalis and E. faecium. Using each of the E. faecalis- and E. faecium-specific PCR primer pairs as well as vanA-, vanB- and vanC-specific PCR primers used alone or in multiplex as described above, the sensitivity ranged from 1 to 10 copies of genomic DNA.

The format of the assay is not limited to the one described above. A person skilled in the art could adapt the assay for different formats such as PCR with real-time detection using molecular beacon probes. Molecular beacon probes designed to be used in this assay include, but are not limited to, SEQ ID NO. 1238 for the detection of *E. faecalis*, SEQ ID NO. 1237 for the detection of *E. faecium*, SEQ ID NO. 1239 for the detection of *vanA*, and SEQ ID NO. 1241 for the detection of *vanB*.

Alternatively, another PCR assay was developed for the detection of vancomycin-resistant *E. faecium* and vancomycin-resistant *E. faecalis*. This assay included two multiplex: (1) the first multiplex contained the *vanA*-specific primer pair (SEQ ID NOs. 1090-1091) and the *vanB*-specific PCR primer pair described in our assigned US patent 5,994,066 (SEQ ID NOs. 1095 and 1096 in the present patent and SEQ ID NOs. 231 and 232 in the said patent), and (2) the second multiplex contained the *E. faecalis*-specific PCR primer pair described in our assigned US patent 5,994,066 (SEQ ID NOs. 40 and 41 in the said patent) and the *E. faecium*-specific PCR primer pair described in our patent publication WO98/20157 (SEQ ID NOs. 1

and 2 in the said publication). For both multiplexes, the optimal cycling conditions for maximum sensitivity and specificity were found to be 3 min. at 94 °C, followed by forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 58 °C, plus a terminal extension at 72 °C for 2 minutes. Detection of the PCR products was made by electrophoresis in agarose gels (2 %) containing 0.25 μ g/ml of ethidium bromide. The two multiplexes were tested for their specificity by using 0.1 nanogram of purified genomic DNA from a panel of two vancomycin-sensitive E. faecalis strains, two vancomycin-resistant E. faecalis strains, two vancomycinsensitive E. faecium strains, two vancomycin-resistant E. faecium strains, 16 other enterococcal species and 31 other gram-positive bacterial species. All the E. faecium and E. faecalis strains were amplified with high specificty showing a perfect correlation between the genotypic analysis and the susceptibility to glycopeptide antibiotics (vancomycin and teicoplanin). The sensitivity of the assay was determined for two vancomycin-resistant E. faecalis strains and two vancomycin-resistant E. faecium strains. The detection limit was 5 copies of genomic DNA for all the strains.

This multiplex PCR assay was coupled with capture-probe hybridization. Four internal probes were designed: one specific to the *vanA* amplicon (SEQ ID NO. 2292), one specific to the *vanB* amplicon (SEQ ID NO. 2294), one specific to the *E. faecalis* amplicon (SEQ ID NO. 2291) and one specific to the *E. faecium* amplicon (SEQ ID NO. 2287). Each of the internal probes detected their specific amplicons with high specificity and sensitivity.

EXAMPLE 24:

Universal amplification involving the EF-G (fusA) subdivision of tuf sequences. As shown in Figure 3, primers SEQ ID NOs. 1228 and 1229 were designed to amplify the region between the end of fusA and the beginning of tuf genes in the str operon. Genomic DNAs from a panel of 35 strains were tested for PCR amplification with those primers. In the initial experiment, the following strains showed a positive

result: Abiotrophia adiacens ATCC 49175, Abiotrophia defectiva ATCC 49176, Bacillus subtilis ATCC 27370, Closridium difficile ATCC 9689, Enterococcus avium ATCC 14025, Enterococcus casseliflavus ATCC 25788, Enterococcus cecorum ATCC 43198, Enterococcus faecalis ATCC 29212, Enterococcus faecium ATCC 19434, Enterococcus flavescens ATCC 49996, Enterococcus gallinarum ATCC 49573, Enterococcus solitarius ATCC 49428, Escherichia coli ATCC 11775, Haemophilus influenzae ATCC 9006, Lactobacillus acidophilus ATCC 4356, Peptococcus niger ATCC 27731, Proteus mirabilis ATCC 25933, Staphylococcus aureus ATCC 43300, Staphylococcus auricularis ATCC 33753, Staphylococcus capitis ATCC 27840, Staphylococcus epidemidis ATCC 14990, Staphylococcus haemolyticus ATCC 29970, Staphylococcus hominis ATCC 27844, Staphylococcus lugdunensis ATCC 43809, Staphylococcus saprophyticus ATCC 15305, Staphylococcus simulans ATCC 27848, and Staphylococcus warneri ATCC 27836. This primer pair could amplify additional bacterial species; however, there was no amplification for some species, suggesting that the PCR cycling conditions could be optimized or the primers modified. For example, SEQ ID NO. 1227 was designed to amplify a broader range of species.

In addition to other possible primer combinations to amplify the region covering fusA and tuf, Figure 3 illustrates the positions of amplification primers SEQ ID NOs. 1221-1227 which could be used for universal amplification of fusA segments. All of the above mentioned primers (SEQ ID NOs. 1221-1229) could be useful for the universal and/or the specific detection of bacteria.

Moreover, different combinations of primers SEQ ID NOs. 1221-1229, sometimes in combination with *tuf* sequencing primer SEQ ID NO. 697, were used to sequence portions of the *str* operon, including the intergenic region. In this manner, the following sequences were generated: SEQ ID NOs. 1518-1526, 1578-1580, 1786-1821, 1822-1834, 1838-1843, 2184, 2187, 2188, 2214-2249, and 2255-2269.

EXAMPLE 25:

DNA fragment isolation from Staphylococcus saprophyticus by arbitrarily primed PCR. DNA sequences of unknown coding potential for the species-specific detection and identification of Staphylococcus saprophyticus were obtained by the method of arbitrarily primed PCR (AP-PCR).

AP-PCR is a method which can be used to generate specific DNA probes for microorganisms (Fani et al., 1993, Molecular Ecology 2:243-250). A description of the AP-PCR protocol used to isolate a species-specific genomic DNA fragment from Staphylococcus saprophyticus follows. Twenty different oligonucleotide primers of 10 nucleotides in length (all included in the AP-PCR kit OPAD (Operon Technologies, Inc., Alameda, CA)) were tested systematically with DNAs from 5 bacterial strains of Staphylococcus saprophyticus as well as with bacterial strains of 27 other staphylococcal (non-S. saprophyticus) species. For all bacterial species, amplification was performed directly from one μL (0.1 ng/ μL) of purified genomic DNA. The 25 µL PCR reaction mixture contained 50 mM KCl, 10 mM Tris-HCl (pH 9.0), 0.1% Triton X-100, 2.5 mM MgCl₂, 1.2 μ M of only one of the 20 different AP-PCR primers OPAD, 200 µM of each of the four dNTPs, 0.5 U of Taq DNA polymerase (Promega Corp., Madison, Wis.) coupled with TaqStartTM antibody (Clontech Laboratories Inc., Palo Alto, CA). PCR reactions were subjected to cycling using a MJ Research PTC-200 thermal cycler as follows: 3 min at 96 °C followed by 42 cycles of 1 min at 94 °C for the denaturation step, 1 min at 31 °C for the annealing step and 2 min at 72 °C for the extension step. A final extension step of 7 min at 72 °C followed the 42 cycles to ensure complete extension of PCR products. Subsequently, twenty microliters of the PCR-amplified mixture were resolved by electrophoresis on a 1.5 % agarose gel containing 0.25 μg/ml of ethidium bromide. The size of the amplification products was estimated by comparison with a 50-bp molecular weight ladder.

Amplification patterns specific for *Staphylococcus saprophyticus* were observed with the AP-PCR primer OPAD-16 (sequence: 5'-AACGGGCGTC-3'). Amplification with this primer consistently showed a band corresponding to a

DNA fragment of approximately 380 bp for all *Staphylococcus saprophyticus* strains tested but not for any of the other staphylococcal species tested.

The band corresponding to the 380 bp amplicon, specific and ubiquitous for *S. saprophyticus* based on AP-PCR, was excised from the agarose gel and purified using the QIAquickTM gel extraction kit (QIAGEN Inc.). The gel-purified DNA fragment was cloned into the T/A cloning site of the pCR 2.1TM plasmid vector (Invitrogen Inc.) using T4 DNA ligase (New England BioLabs). Recombinant plasmids were transformed into *E. coli* DH5α competent cells using standard procedures. All reactions were performed according to the manufacturer's instructions. Plasmid DNA isolation was done by the method of Birnboim and Doly (Nucleic Acid Res., 1979, 7:1513-1523) for small-scale preparations. All plasmid DNA preparations were digested with the EcoRI restriction endonuclease to ensure the presence of the approximately 380 bp AP-PCR insert into the plasmid. Subsequently, a large-scale and highly purified plasmid DNA preparation was performed from two selected clones shown to carry the AP-PCR insert by using the QIAGEN plasmid purification kit (midi format). These large-scale plasmid preparations were used for automated DNA sequencing.

The 380 bp nucleotide sequence was determined for three strains of *S. saprophyticus* (SEQ ID NOs. 74, 1093, and 1198). Both strands of the AP-PCR insert from the two selected clones were sequenced by the dideoxynucleotide chain termination sequencing method with SP6 and T7 sequencing primers by using the Applied Biosystems automated DNA sequencer (model 373A) with their PRISMTM Sequenase^{RTM} Terminator Double-stranded DNA Sequencing Kit (Applied Biosystems, Foster City, CA).

Optimal species-specific amplification primers (SEQ ID NOs. 1208 and 1209) have been selected from the sequenced AP-PCR Staphylococcus saprophyticus DNA fragments with the help of the primer analysis software OligoTM 5.0 (National BioSciences Inc.). The selected primers were tested in PCR assays to verify their specificity and ubiquity. Data obtained with DNA preparations from reference ATCC strains of 49 gram-positive and 31 gram-negative bacterial

species, including 28 different staphylococcal species, indicate that the selected primer pairs are specific for *Staphylococcus saprophyticus* since no amplification signal has been observed with DNAs from the other staphylococcal or bacterial species tested. This assay was able to amplify efficiently DNA from all 60 strains of *S. saprophyticus* from various origins tested. The sensitivity level achieved for three *S. saprophyticus* reference ATCC strains was around 6 genome copies.

EXAMPLE 26:

Sequencing of prokaryotic *tuf* gene fragments. The comparison of publicly available *tuf* sequences from a variety of bacterial species revealed conserved regions, allowing the design of PCR primers able to amplify *tuf* sequences from a wide range of bacterial species. Using primer pair SEQ ID NOs. 664 and 697, it was possible to amplify and determine *tuf* sequences SEQ ID NOs.: 1-73, 75-241, 607-618, 621, 662, 675, 717-736, 868-888, 932, 967-989, 992, 1002, 1572-1575, 1662-1663, 1715-1733, 1835-1837, 1877-1878, 1880-1881, 2183, 2185, 2200, 2201, and 2270-2272.

EXAMPLE 27:

Sequencing of procaryotic recA gene fragments. The comparison of publicly available recA sequences from a variety of bacterial species revealed conserved regions, allowing the design of PCR primers able to amplify recA sequences from a wide range of bacterial species. Using primer pairs SEQ ID NOs. 921-922 and 1605-1606, it was possible to amplify and determine recA sequences SEQ ID NOs.: 990-991, 1003, 1288-1289, 1714, 1756-1763, 1866-1873 and 2202-2212.

EXAMPLE 28:

Specific detection and identification of Escherichia coli/Shigella sp. using tuf sequences. The analysis of tuf sequences from a variety of bacterial species allowed the selection of PCR primers (SEQ ID NOs. 1661 and 1665) and of an internal probe (SEQ ID NO. 2168) specific to Escherichia coli/Shigella sp. The strategy used to design the PCR primers was based on the analysis of a multiple sequence alignment of various tuf sequences. The multiple sequence alignment included the tuf sequences of Escherichia coli/Shigella sp. as well as tuf sequences from other species and bacterial genera, especially representatives of closely related species. A careful analysis of this alignment allowed the selection of oligonucleotide sequences which are conserved within the target species but which discriminate sequences from other species, especially from the closely related species, thereby permitting the species-specific and ubiquitous detection and identification of the target bacterial species.

The chosen primer pair, oligos SEQ ID NOs. 1661 and 1665, gives an amplification product of 219 bp. Standard PCR was carried out using 0.4 μ M of each primer, 2.5 mM MgCl₂, BSA 0.05 mM, 50 mM KCl, 10 mM Tris-HCl (pH 9.0), 0.1 % Triton X-100, dNTPs 0.2 mM (Pharmacia), 0,5 U *Taq* DNA polymerase (Promega) coupled with TaqStartTM antibody (Clontech Laboratories Inc.), 1 μ l of genomic DNA sample in a final volume of 20 μ l using a PTC-200 thermocycler (MJ Research). The optimal cycling conditions for maximum sensitivity and specificity were 3 minutes at 95 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 60 °C, followed by terminal extension at 72 °C for 2 minutes. Detection of the PCR products was made by electrophoresis in agarose gels (2 %) containing 0.25 μ g/ml of ethidium bromide. Visualization of the PCR products was made under UV at 254 nm.

Specificity of the assay was tested by adding to the PCR reactions 0.1 ng of genomic DNA from each of the following bacterial species: Escherichia coli (7

strains), Shigella sonnei, Shigella flexneri, Shigella dysenteriae, Salmonella typhimyurium, Salmonella typhi, Salmonella enteritidis, Tatumella ptyseos, Klebsiella pneumoniae (2 strains), Enterobacter aerogenes, Citrobacter farmeri, Campylobacter jejuni, Serratia marcescens. Amplification was observed only for the Escherichia coli and Shigella sp. strains listed and Escherichia fergusonii. The sensitivity of the assay with 40-cycle PCR was verified with one strain of E. coli and three strains of Shigella sp. The detection limit for E. coli and Shigella sp. was 1 to 10 copies of genomic DNA, depending on the strains tested.

EXAMPLE 29:

Specific detection and identification of *Klebsiella pneumoniae* using *atpD* sequences. The analysis of *atpD* sequences from a variety of bacterial species allowed the selection of PCR primers specific to *K. pneumoniae*. The primer design strategy is similar to the strategy described in Example 28 except that *atpD* sequences were used in the alignment.

Two K. pneumoniae-specific primers were selected, (SEQ ID NOs. 1331 and 1332) which give an amplification product of 115 bp. Standard PCR was carried out on PTC-200 thermocyclers (MJ Research) using 0.4 μ M of each primer as described in Example 28. The optimal cycling conditions for maximum sensitivity and specificity were as follow: three minutes at 95 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 55°C, followed by terminal extension at 72 °C for 2 minutes.

Specificity of the assay was tested by adding to the PCR reactions 0.1 ng of genomic DNA from each of the following bacterial species: Klebsiella pneumoniae (2 strains), Klebsiella ornitholytica, Klebsiella oxytoca (2 strains), Klebsiella planticola, Klebsiella terrigena, Citrobacter freundii, Escherichia coli, Salmonella cholerasuis typhi, Serratia marcescens, Enterobacter aerogenes, Proteus vulgaris,

Kluyvera ascorbata, Kluyvera georgiana, Kluyvera cryocrescens and Yersinia enterolitica. Amplification was detected for the two K. pneumoniae strains, K. planticola, K. terrigena and the three Kluyvera species tested. Analysis of the multiple alignment sequence of the atpD gene allowed the design of an internal probe SEQ ID NO. 2167 which can discrimate Klebsiella pneumoniae from other Klebsiella sp. and Kluyvera sp. The sensitivity of the assay with 40-cycle PCR was verified with one strain of K. pneumoniae. The detection limit for K. pneumoniae was around 10 copies of genomic DNA.

EXAMPLE 30:

Specific detection and identification of Acinetobacter baumannii using atpD sequences. The analysis of atpD sequences from a variety of bacterial species allowed the selection of PCR primers specific to Acinetobacter baumannii. The primer design strategy is similar to the strategy described in Example 28.

Two A. baumannii-specific primers were selected, SEQ ID NOs. 1690 and 1691, which give an amplification product of 233 bp. Standard PCR was carried out on PTC-200 thermocyclers (MJ Research) using 0.4 μ M of each primer as described in Example 28. The optimal cycling conditions for maximum sensitivity and specificity were as follow: three minutes at 95 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 60°C, followed by terminal extension at 72 °C for 2 minutes.

Specificity of the assay was tested by adding to the PCR reactions 0.1 ng of genomic DNA from each of the following bacterial species: Acinetobacter baumannii (3 strains), Acinetobacter anitratus, Acinetobacter lwoffi, Serratia marcescens, Enterobacter cloacae, Enterococcus faecalis, Pseudomonas aeruginosa, Psychrobacter phenylpyruvicus, Neisseria gonorrheoae, Haemophilus haemoliticus, Yersinia enterolitica, Proteus vulgaris, Eikenella corrodens,

Escherichia coli. Amplification was detected only for A. baumannii, A anitratus and A. lwoffi. The sensitivity of the assay with 40-cycle PCR was verified with two strains of A. baumannii. The detection limit for the two A. baumannii strains tested was 5 copies of genomic DNA. Analysis of the multiple alignment sequence of the atpD gene allowed the design of a A. baumannii-specific internal probe (SEQ ID NO. 2169).

EXAMPLE 31:

Specific detection and identification of *Neisseria gonorrhoeae* using *tuf* sequences. The analysis of *tuf* sequences from a variety of bacterial species allowed the selection of PCR primers specific to *Neisseria gonorrhoeae*. The primer design strategy is similar to the strategy described in Example 28.

Two N. gonorrhoeae-specific primers were selected, SEQ ID NOs. 551 and 552, which give an amplification product of 139 bp. PCR amplification was carried out on PTC-200 thermocyclers (MJ Research) using 0.4 μ M of each primer as described in Example 28. The optimal cycling conditions for maximum sensitivity and specificity were as follow: three minutes at 95 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 65°C, followed by terminal extension at 72 °C for 2 minutes.

Specificity of the assay was tested by adding into the PCR reactions, 0.1 ng of genomic DNA from each of the following bacterial species: Neisseria gonorrhoeae (19 strains), Neisseria meningitidis (2 strains), Neisseria lactamica, Neisseria flavescens, Neisseria animalis, Neisseria canis, Neisseria cuniculi, Neisseria elongata, Neisseria mucosa, Neisseria polysaccharea, Neisseria sicca, Neisseria subflava, Neisseria weaveri. Amplification was detected only for N. gonorrhoeae, N. sicca and N. polysaccharea. The sensitivity of the assay with 40-cycle PCR was verified with two strains of N. gonorrhoeae. The detection limit for the N.

gonorrhoeae strains tested was 5 copies of genomic DNA. Analysis of the multiple alignment sequence of the *tuf* gene allowed the design of an internal probe, SEQ ID NO. 2166, which can discriminate N. gonorrhoeae from N. sicca and N. polysaccharea.

EXAMPLE 32:

Sequencing of bacterial gyrA and parC gene fragments. Sequencing of bacterial gyrA and parC fragments. One of the major mechanism of resistance to quinolone in various bacterial species is mediated by target changes (DNA gyrase and/or topoisomerase IV). These enzymes control DNA topology and are vital for chromosome function and replication. Each of these enzymes is a tetramer composed of two subunits: GyrA and GyrB forming A₂B₂ complex in DNA gyrase; and ParC and ParE forming and C₂E₂ complex in DNA topoisomerase IV. It has been shown that they are hotspots, called the quinolone-resitance-determining region (QRDR) for mutations within gyrA that encodes for the GyrA subunit of DNA gyrase and within parC that encodes the parC subunit of topoisomerase IV.

In order to generate a database for gyrA and parC sequences that can be used for design of primers and/or probes for the specific detection of quinolone resistance in various bacterial species, gyrA and parC DNA fragments selected from public database (GenBanK and EMBL) from a variety of bacterial species were used to design oligonucleotide primers.

Using primer pair SEQ ID NOs. 1297 and 1298, it was possible to amplify and determine gyrA sequences from Klebsiella oxytoca (SEQ ID NO. 1764), Klebsiella pneumoniae subsp. ozaneae (SEQ ID NO. 1765), Klebsiella planticola (SEQ ID NO. 1766), Klebsiella pneumoniae (SEQ ID NO. 1767), Klebsiella pneumoniae subsp. pneumoniae (two strains) (SEQ ID NOs. 1768-1769), Klebsiella

pneumoniae subsp. rhinoscleromatis (SEQ ID NO. 1770), Klebsiella terrigena. (SEQ ID NO. 1771), Kluyvera ascorbata (SEQ ID NO. 2013), Kluyvera georgiana (SEQ ID NO. 2014) and Escherichia coli (4 strains) (SEQ ID NOs. 2277-2280). Using primer pair SEQ ID NOs. 1291 and 1292, it was possible to amplify and determine gyrA sequences from Legionella pneumophila subsp. pneumophila (SEQ ID NO. 1772), Proteus mirabilis (SEQ ID NO. 1773), Providencia rettgeri (SEQ ID NO. 1774), Proteus vulgaris (SEQ ID NO. 1775) and Yersinia enterolitica (SEQ ID NO. 1776). Using primer pair SEQ ID NOs. 1340 and 1341, it was possible to amplify and determine gyrA sequence from Staphylococcus aureus (SEQ ID NO. 1255).

Using primers SEQ ID NOs. 1318 and 1319, it was possible to amplify and determine parC sequences from K. oxytoca (two strains) (SEQ ID NOs. 1777-1778), Klebsiella pneumoniae subsp. ozaenae (SEQ ID NO. 1779), Klebsiella planticola (SEQ ID NO. 1780), Klebsiella pneumoniae (SEQ ID NO. 1781), Klebsiella pneumoniae subsp. pneumoniae (two strains) (SEQ ID NOs. 1782-1783), Klebsiella pneumoniae subsp. rhinoscleromatis (SEQ ID NO. 1784) and Klebsiella terrigena (SEQ ID NO. 1785).

EXAMPLE 33:

Development of a PCR assay for the specific detection and identification of Staphylococcus aureus and its quinolone resistance genes gyrA and parC. The analysis of gyrA and parC sequences from a variety of bacterial species revealed conserved regions allowing the design of PCR primers specific to the quinolone-resistance-determining region (QRDR) of gyrA and parC from Staphylococcus aureus. PCR primer pair SEQ ID NOs. 1340 and 1341 was designed to amplify the gyrA sequence of S. aureus, whereas PCR primer pair SEQ ID NOs. 1342 and 1343 was designed to amplify S. aureus parC. The comparison of gyrA and parC sequences from S. aureus strains with various levels of quinolone resistance

allowed the identification of amino acid substitutions Ser-84 to Leu, Glu-88 to Gly or Lys in the GyrA subunit of DNA gyrase encoded by gyrA and amino acid changes Ser-80 to Phe or Tyr and Ala-116 to Glu in the ParC subunit of topoisomerase IV encoded by parC. These amino acid substitutions in GyrA and ParC subunits occur in isolates with intermediate- or high-level quinolone resistance. Internal probes for the specific detection of wild-type S. aureus gyrA (SEQ ID NO. 1940) and wild-type S. aureus parC (SEQ ID NO. 1941) as well as internal probes for the specific detection of each of the gyrA (SEQ ID NOs. 1333-1335) and parC mutations identified in quinolone-resistant S. aureus (SEQ ID NOs. 1336-1339) were designed.

The gyrA- and parC-specific primer pairs (SEQ ID NOs. 1340-1341 and SEQ ID NOs. 1342-1343) were used in multiplex. PCR amplification was carried out on PTC-200 thermocyclers (MJ Research) using 0.3, 0.3, 0.6 and 0.6 μM of each primers, respectively, as described in Example 28. The optimal cycling conditions for maximum sensitivity and specificity were 3 minutes at 95 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 62 °C, followed by terminal extension at 72 °C for 2 minutes. Detection of the PCR products was made by electrophoresis in agarose gels (2 %) containing $0.25 \mu \text{g/ml}$ of ethidium bromide. The specificity of the multiplex assay with 40cycle PCR was verified by using 0.1 ng of purified genomic DNA from a panel of gram-positive bacteria. The list included the following: Abiotrophia adiacens, Abiotrophia defectiva, Bacillus cereus, Bacillus mycoides, Enterococcus faecalis (2 strains), Enterococcus flavescens, Gemella morbillorum, Lactococcus lactis, Listeria innocua, Listeria monocytogenes, Staphylococcus aureus (5 strains), capitis subsp. urealyticus, Staphylococcus Staphylococcus auricalis, Staphylococcus Staphylococcus chromogenes, Staphylococcus carnosus, epidermidis (3 strains), Staphylococcus gallinarum, Staphylococcus haemolyticus (2 strains), Staphylococcus hominis, Staphylococcus hominis subsp hominis, Staphylococcus Staphylococcus lugdunensis, Staphylococcuslentus,

saccharolyticus, Staphylococcus saprophyticus (3 strains), Staphylococcus simulans, Staphylococcus warneri, Staphylococcus xylosus, Streptococcus agalactiae, Streptococcus pneumoniae. Strong amplification of both gyrA and parC genes was only detected for the S. aureus strains tested. The sensitivity of the multiplex assay with 40-cycle PCR was verified with one quinolone-sensitive and four quinolone-resistant strains of S. aureus. The detection limit was 2 to 10 copies of genomic DNA, depending on the strains tested.

Detection of the hybridization with the internal probes was performed as described in Example 7. The internal probes specific to wild-type gyrA and parC of S. aureus and to the gyrA and parC variants of S. aureus were able to recognize two quinolone-resistant and one quinolone-sensitive S. aureus strains showing a perfect correlation with the susceptibility to quinolones.

The complete assay for the specific detection of *S. aureus* and its susceptibility to quinolone contains the *Staphylococcus*-specific primers (SEQ ID NOs. 553 and 575) described in Example 7 and the multiplex containing the *S. aureus gyrA*- and parC-specific primer pairs (SEQ ID NOs. 1340-1341 and SEQ ID NOs. 1342-1343). Amplification is coupled with post-PCR hybridization with the internal probe specific to *S. aureus* (SEQ ID NO. 587) described in Example 7 and the internal probes specific to wild-type *S. aureus gyrA* and parC (SEQ ID NOs. 1940-1941) and to the *S. aureus gyrA* and parC variants (SEQ ID NOs. 1333-1338).

An assay was also developed for the detection of quinolone-resistant *S. aureus* using the SmartCycler (Cepheid). Real-time detection is based on the use of *S. aureus parC*-specific primers (SEQ ID NOs. 1342 and 1343) and the *Staphylococcus*-specific primers (SEQ ID NOs. 553 and 575) described in Example 7. Internal probes were designed for molecular beacon detection of the wild-type *S. aureus parC* (SEQ ID NO.1939), for detection of the Ser-80 to Tyr or

Phe amino acid substitutions in the ParC subunit encoded by S. aureus parC (SEQ ID NOs. 1938 and 1955) and for detection of S. aureus (SEQ ID NO. 2282).

EXAMPLE 34:

Development of a PCR assay for the detection and identification of Klebsiella pneumoniae and its quinolone resistance genes gyrA and parC. The analysis of gyrA and parC sequences from a variety of bacterial species from the public databases and from the database described in Example 32 revealed conserved regions allowing the design of PCR primers specific to the quinolone-resistancedetermining region (QRDR) of gyrA and parC from K. pneumoniae. PCR primer pair SEQ ID NOs. 1936 and 1937, or pair SEQ ID NOs. 1937 and 1942, were designed to amplify the gyrA sequence of K. pneumoniae, whereas PCR primer pair SEQ ID NOs. 1934 and 1935 was designed to amplify K. pneumoniae parC sequence. An alternative pair, SEQ ID NOs. 1935 and 1936, can also amplify K. pneumoniae parC. The comparison of gyrA and parC sequences from K. pneumoniae strains with various levels of quinolone resistance allowed the identification of amino acid substitutions Ser-83 to Tyr or Phe and Asp-87 to Gly or Ala and Asp-87 to Asn in the GyrA subunit of DNA gyrase encoded by gyrA and amino acid changes Ser-80 to Ile or Arg and Glu-84 to Gly or Lys in the ParC subunit of topoisomerase IV encoded by parC. These amino acid substitutions in the GyrA and ParC subunits occur in isolates with intermediate- or high-level quinolone resistance. Internal probes for the specific detection of wild-type K. pneumoniae gyrA (SEQ ID NO. 1943) and wild-type K. pneumoniae parC (SEQ ID NO. 1944) as well as internal probes for the specific detection of each of the gyrA (SEQ ID NOs. 1945-1949) and parC mutations identified in quinoloneresistant K. pneumoniae (SEQ ID NOs. 1950-1953) were designed.

Two multiplex using the K. pneumoniae gyrA- and parC-specific primer pairs were used: the first multiplex contained K. pneumoniae gyrA-specific primers (SEQ ID

NOs. 1937 and 1942) and K. pneumoniae parC-specific primers (SEQ ID NOs. 1934 and 1935) and the second multiplex contained K. pneumoniae gyrA/parCspecific primer (SEQ ID NOs. 1936), K. pneumoniae gyrA-specific primer (SEQ ID NO. 1937) and K. pneumoniae parC-specific primer (SEQ ID NO. 1935). Standard PCR was carried out on PTC-200 thermocyclers (MJ Research) using for the first multiplex 0.6, 0.6, 0.4, 0.4 μM of each primer, respectively, and for the second multiplex 0.8, 0.4, 0.4 μM of each primer, respectively. PCR amplification and agarose gel electrophoresis of the amplified products were performed as described in Example 28. The optimal cycling conditions for maximum sensitivity and specificity were 3 minutes at 95 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 62 °C, followed by terminal extension at 72 °C for 2 minutes. The specificity of the two multiplex assays with 40-cycle PCR was verified by using 0.1 ng of purified genomic DNA from a panel of gram-negative bacteria. The list included: Acinetobacter baumannii, Citrobacter freundii, Eikenella corrodens, Enterobacter aerogenes, Enterobacter cancerogenes, Enterobacter cloacae, Escherichia coli (10 strains), Haemophilus influenzae, Klebsiella pneumoniae, Klebsiella ornitholytica, Klebsiella oxytoca (2 strains), Klebsiella planticola, Klebsiella terrigena, Kluyvera ascorbata, Kluyvera cryocrescens, Kluyvera georgiana, Neisseria gonorrhoeae, Proteus mirabilis, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella choleraesuis subsp. typhimurium, Salmonella enteritidis, Serratia liquefaciens, Serratia marcescens and Yersinia enterocolytica. For both multiplex, strong amplification of both gyrA and parC was observed only for the K. pneumoniae strain tested. The sensitivity of the two multiplex assays with 40-cycle PCR was verified with one quinolone-sensitive strain of K. pneumoniae. The detection limit was around 10 copies of genomic DNA.

The complete assay for the specific detection of *K. pneumoniae* and its susceptibility to quinolone contains the *Klebsiella*-specific primers (SEQ ID NOs. 1331 and 1332) described in Example 29 and either the multiplex containing the *K*.

pneumoniae gyrA- and parC-specific primers (SEQ ID NOs. 1935, 1936, 1937) or the multiplex containing the K. pneumoniae gyrA- and parC-specific primers (SEQ ID NOs. 1934, 1937, 1939, 1942). Amplification is coupled with post-PCR hybridization with the internal probe specific to K. pneumoniae (SEQ ID NO. 2167) described in Example 29 and the internal probes specific to wild-type K. pneumoniae gyrA and parC (SEQ ID NOs. 1943, 1944) and to the K. pneumoniae gyrA and parC variants (SEQ ID NOs. 1945-1949 and 1950-1953).

An assay was also developed for the detection of quinolone-resistant *K. pneumoniae* using the SmartCycler (Cepheid). Real-time detection is based on the use of resistant *K. pneumoniae gyrA*-specific primers (SEQ ID NOs. 1936 and 1937) and the *K. pneumoniae*-specific primers (SEQ ID NOs. 1331 and 1332) described in Example 29. Internal probes were designed for molecular beacon detection of the wild-type *K. pneumoniae gyrA* (SEQ ID NO. 2251), for detection of the Ser-83 to Tyr or Phe and/or Asp-87 to Gly or Asn in the GyrA subunit of DNA gyrase encoded by *gyrA* (SEQ ID NOs. 2250) and for detection of *K. pneumoniae* (SEQ ID NO. 2281).

EXAMPLE 35:

Development of a PCR assay for detection and identification of S. pneumoniae and its quinolone resistance genes gyrA and parC. The analysis of gyrA and parC sequences from a variety of bacterial species revealed conserved regions allowing the design of PCR primers able to amplify the quinolone-resistance-determining region (QRDR) of gyrA and parC from all S. pneumoniae strains. PCR primer pair SEQ ID NOs. 2040 and 2041 was designed to amplify the QRDR of S. pneumoniae gyrA, whereas PCR primer pair SEQ ID NOs. 2044 and 2045 was designed to amplify the QRDR of S. pneumoniae parC. The comparison of gyrA and parC sequences from S. pneumoniae strains with various levels of quinolone resistance allowed the identification of amino acid substitutions Ser-81 to Phe or

Tyr in the GyrA subunit of DNA gyrase encoded by gyrA and amino acid changes Ser-79 to Phe in the ParC subunit of topoisomerase IV encoded by parC. These amino acid substitutions in the GyrA and ParC subunits occur in isolates with intermediate- or high-level quinolone resistance. Internal probes for the specific detection of each of the gyrA (SEQ ID NOs. 2042 and 2043) and parC (SEQ ID NO. 2046) mutations identified in quinolone-resistant S. pneumoniae were designed.

For all bacterial species, amplification was performed from purified genomic DNA. 1 µl of genomic DNA at 0.1 ng/µL was transferred directly to a 19 µl PCR mixture. Each PCR reaction contained 50 mM KCl, 10 mM Tris-HCl (pH 9.0), 0.1% Triton X-100, 2.5 mM MgCl₂, 0.4 µM (each) of the above primers SEQ ID NOs. 2040, 2041, 2044 and 2045, 0.05 mM bovine serum albumin (BSA) and 0.5 U *Taq* polymerase coupled with TaqStartTM antibody. The optimal cycling conditions for maximum sensitivity and specificity were 3 minutes at 95 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 58 °C, followed by terminal extension at 72 °C for 2 minutes. In order to generate Digoxigenin (DIG)-labeled amplicons for capture probe hybridization, 0.1X PCR DIG labeling four deoxynucleoside triphosphates mix (Boehringer Mannheim GmbH) was used for amplification.

The DIG-labeled amplicons were hybridized to the capture probes bound to 96-well plates. The plates were incubated with anti-DIG-alkaline phosphatase and the chemiluminescence was measured by using a luminometer (MLX, Dynex Technologies Inc.) after incubation with CSPD and recorded as Relative Light Unit (RLU). The RLU ratio of tested sample with and without captures probes was then calculated. A ratio ≥ 2.0 was defined as a positive hybridization signal. All reactions were performed in duplicate.

The specificity of the multiplex assay with 40-cycle PCR was verified by using 0.1 ng of purified genomic DNA from a panel of bacteria listed in Table 13. Strong amplification of both gyrA and parC was detected only for the S. pneumoniae strains tested. Weak amplification of both gyrA and parC genes was detected for Staphylococcus simulans. The detection limit tested with purified genomic DNA from 5 strains of S. pneumoniae was 1 to 10 genome copies. In addition, 5 quinolone-resistant and 2 quinolone-sensitive clinical isolates of S. pneumoniae were tested to further validate the developed multiplex PCR coupled with capture probe hybridization assays. There was a perfect correlation between detection of S. pneumoniae gyrA and parC mutations and the susceptibility to quinolone.

The complete assay for the specific detection of *S. pneumoniae* and its susceptibility to quinolone contains the *S. pneumoniae*-specific primers (SEQ ID NOs. 1179 and 1181) described in Exemple 20 and the multiplex containing the *S. pneumoniae gyrA*-specific and *parC*-specific primer pairs (SEQ ID NOS. 2040 and 2041 and SEQ ID NOs. 2044 and 2045). Amplification is coupled with post-PCR hybridization with the internal probe specific to *S. pneumoniae* (SEQ ID NO. 1180) described in Example and the internal probes specific to each of the *S. pneumoniae gyrA* and *parC* variants (SEQ ID NOs. 2042, 2043 and 2046).

EXAMPLE 36:

Detection of extended-spectrum TEM-type β-lactamases in *Escherichia coli*. The analysis of TEM sequences which confer resistance to third-generation cephalosporins and to β-lactamase inhibitors allowed the identification of amino acid substitutions Met-69 to Ile or Leu or Val, Ser-130 to Gly, Arg-164 to Ser or His, Gly-238 to Ser, Glu-240 to Lys and Arg-244 to Ser or Cys or Thr or His or Leu. PCR primers SEQ ID NOs. 1907 and 1908 were designed to amplify TEM sequences. Internal probes for the specific detection of wild-type TEM (SEQ ID NO. 2141) and for each of the amino acid substitutions (SEQ ID NOs. 1909-1926) identified in TEM variants were designed to detect resistance to third-generation

cephalosporins and to β -lactamase inhibitors. Design and synthesis of primers and probes, and detection of the hybridization were performed as described in Example 7.

For all bacterial species, amplification was performed from purified genomic DNA. One μ l of genomic DNA at 0.1ng/ μ l was transferred directly to a 19 μ l PCR mixture. Each PCR reaction contained 50 mM KCl, 10 mM Tris-HCl (pH 9.0); 0.1% Triton X-100, 2.5 mM MgCl₂, 0.4 μ M of the TEM-specific primers SEQ ID NOs. 1907 and 1908, 200 μ M (each) of the four deoxynucleoside triphosphates, 0.05 mM bovine serum albumin (BSA) and 0.5 U *Taq* polymerase (Promega) coupled with TaqStartTM antibody. PCR amplification and agarose gel analysis of the amplified products were performed as described in Example 28. The optimal cycling conditions for maximum sensitivity and specificity were 3 minutes at 95 °C for initial denaturation, then forty cycles of three steps consisting of 5 seconds at 95 °C, 30 seconds at 55 °C and 30 seconds at 72 °C, followed by terminal extension at 72 °C for 2 minutes.

The specificity of the TEM-specific primers with 40-cycle PCR was verified by using 0.1 ng of purified genomic from the following bacteria: three third-generation cephalosporin-resistant *Escherichia coli* strains (one with TEM-10, one with TEM-28 and the other with TEM-49), two third-generation cephalosporin-sensitive *Escherichia coli* strain (one with TEM-1 and the other without TEM), one third-generation cephalosporin-resistant *Klebsiella pneumoniae* strain (with TEM-47), and one β -lactamase-inhibitor-resistant *Proteus mirabilis* strain (with TEM-39). Amplification with the TEM-specific primers was detected only for strains containing TEM.

The sensitivity of the assay with 40-cycle PCR was verified with three E. coli strains containing TEM-1 or TEM-10 or TEM-49, one K. pneumoniae strain containing TEM-47 and one P. mirabilis strain containing TEM-39. The detection

limit was 5 to 100 copies of genomic DNA, depending on the TEM-containing strains tested.

The TEM-specific primers SEQ ID NOs. 1907 and 1908 were used in multiplex with the *Escherichia coli/Shigella sp.*-specific primers SEQ ID NOs. 1661 and 1665 described in Example 28 to allow the complete identification of *Escherichia coli/Shigella sp.* and the susceptibility to β -lactams. PCR amplification with 0.4 μ M of each of the primers and agarose gel analysis of the amplified products was performed as described above.

The specificity of the multiplex with 40-cycle PCR was verified by using 0.1 ng of purified genomic DNA from the following bacteria: three third-generation cephalosporin-resistant *Escherichia coli* strains (one with TEM-10, one with TEM-28 and the other with TEM-49), two third-generation cephalosporin-sensitive *Escherichia coli* strain (one with TEM-1 and the other without TEM), one third-generation cephalosporin-resistant *Klebsiella pneumoniae* strain (with TEM-47), and one β-lactamase-inhibitor-resistant *Proteus mirabilis* strain (with TEM-39). The multiplex was highly specific to *Escherichia coli* strains containing TEM.

The complete assay for detection of TEM-type β-lactamases in *E. coli* includes PCR amplification using the multiplex containing the TEM-specific primers (SEQ ID NOs. 1907 and 1908) and the *Escherichia coli/Shigella* sp.-specific primers (SEQ ID NOs. 1661 and 1665) coupled with post PCR-hybridization with the internal probes specific to wild-type TEM (SEQ ID NO. 2141) and to the TEM variants (SEQ ID NOs. 1909-1926).

EXAMPLE 37:

Detection of extended-spectrum SHV-type β-lactamases in Klebsiella pneumoniae.

The comparison of SHV sequences, which confer resistance to third-generation

cephalosporins and to β -lactamase inhibitors, allowed the identification of amino acid substitutions Ser-130 to Gly, Asp-179 to Ala or Asn, Gly-238 to Ser , and Glu-240 to Lys. PCR primer pair SEQ ID NOs. 1884 and 1885 was designed to amplify SHV sequences. Internal probes for the specific identification of wild-type SHV (SEQ ID NO. 1896) and for each of the amino acid substitutions (SEQ ID NOs. 1886-1895 and 1897-1898) identified in SHV variants were designed to detect resistance to third-generation cephalosporins and to β -lactamase inhibitors. Design and synthesis of primers and probes, and detection of the hybridization were performed as described in Example 7.

For all bacterial species, amplification was performed from purified genomic DNA. One μl of of genomic DNA at 0.1ng/μl was transferred directly to a 19 μl PCR mixture. Each PCR reaction contained 50 mM KCl, 10 mM Tris-HCl (pH 9.0), 0.1% Triton X-100, 2.5 mM MgCl₂, 0.4 μM of the SHV-specific primers SEQ ID NO. 1884 and 1885, 200 μM (each) of the four deoxynucleoside triphosphates, 0.05 mM bovine serum albumin (BSA) and 0.5 U *Taq* polymerase (Promega) coupled with TaqStartTM antibody. PCR amplification and agarose gel analysis of the amplified products were performed as described in Example 28. The optimal cycling conditions for maximum sensitivity and specificity were 3 minutes at 95 °C for initial denaturation, then forty cycles of three steps consisting of 5 seconds at 95 °C, 30 seconds at 55 °C and 30 seconds at 72 °C, followed by terminal extension at 72 °C for 2 minutes.

The specificity of the SHV-specific primers with 40-cycle PCR was verified by using 0.1 ng of purified genomic from the following bacteria: two third-generation cephalosporin-resistant *Klebsiella pneumoniae* strains (one with SHV-2a and the other with SHV-12), one third-generation cephalosporin-sensitive *Klebsiella pneumoniae* strain (with SHV-1), two third-generation cephalosporin-resistant *Escherichia coli* strains (one with SHV-8 and the other with SHV-7), and two third-generation cephalosporin-sensitive *Escherichia coli* strains (one with SHV-1

and the other without any SHV). Amplification with the SHV-specific primers was detected only for strains containing SHV.

The sensitivity of the assay with 40-cycle PCR was verified with four strains containing SHV. The detection limit was 10 to 100 copies of genomic DNA, depending on the SHV-containing strains tested.

The amplification was coupled with post-PCR hybridization with the internal probes specific for identification of wild-type SHV (SEQ ID NO. 1896) and for each of the amino acid substitutions (SEQ ID NOs. 1886-1895 and 1897-1898) identified in SHV variants. The specificity of the probes was verified with six strains containing various SHV enzymes, one *Klebsiella pneumoniae* strain containing SHV-1, one *Klebsiella pneumoniae* strain containing SHV-2a, one *Klebsiella pneumoniae* strain containing SHV-1, one *Escherichia coli* strain containing SHV-7 and one *Escherichia coli* strain containing SHV-8. The probes correctly detected each of the SHV genes and their specific mutations. There was a perfect correlation between the SHV genotype of the strains and the susceptibility to β-lactam antibiotics.

The SHV-specific primers SEQ ID NOs. 1884 and 1885 were used in multiplex with the K. pneumoniae-specific primers SEQ ID NOs. 1331 and 1332 described in Example 29 to allow the complete identification of K. pneumoniae and the susceptibility to β -lactams. PCR amplification with 0.4 μ M of each of the primers and agarose gel analysis of the amplified products were performed as described above.

The specificity of the multiplex with 40-cycle PCR was verified by using 0.1 ng of purified genomic DNA from the following bacteria: three *K. pneumoniae* strains containing SHV-1, one *Klebsiella pneumoniae* strain containing SHV-2a, one

Klebsiella pneumoniae strain containing SHV-12, one K. rhinoscleromatis strain containing SHV-1, one Escherichia coli strain without SHV. The multiplex was highly specific to Klebsiella pneumoniae strain containing SHV.

EXAMPLE 38:

Development of a PCR assay for the detection and identification of *Neisseria* gonorrhoeae and its associated tetracycline resistance gene tetM. The analysis of publicly available tetM sequences revealed conserved regions allowing the design of PCR primers specific to tetM sequences. The PCR primer pair SEQ ID NOs. 1588 and 1589 was used in multiplex with the Neisseria gonorrhoeae-specific primers SEQ ID NOs. 551 and 552 described in Example 31. Sequence alignment analysis of tetM sequences revealed regions suitable for the design of an internal probe specific to tetM (SEQ ID NO. 2254). PCR amplification was carried out on PTC-200 thermocyclers (MJ Research) using 0.4 μ M of each primer pair as described in Example 28. The optimal cycling conditions for maximum sensitivity and specificity were as follow: three minutes at 95 °C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95 °C and 30 seconds at 60°C, followed by terminal extension at 72 °C for 2 minutes.

The specificity of the multiplex PCR assay with 40-cycle PCR was verified by using 0.1 ng of purified genomic DNA from the following bacteria: two tetracycline-resistant *Escherichia coli* strains (one containing the tetracycline-resistant gene *tetB* and the other containing the tetracycline-resistant gene *tetC*), one tetracycline-resistant *Pseudomonas aeruginosa* strain (containing the tetracycline-resistant gene *tetA*), nine tetracycline-resistant *Neisseria gonorrhoeae* strains, two tetracycline-sensitive *Neisseria meningitidis* strains, one tetracycline-sensitive *Neisseria polysaccharea* strain, one tetracycline-sensitive *Neisseria sicca* strain and one tetracycline-sensitive *Neisseria subflava* strain. Amplification with both the *tetM*-specific and *Neisseria gonorrhoeae*-specific primers was detected

only for N. gonorrhoeae strains containing tetM. There was a weak amplification signal using Neisseria gonorrhoeae-specific primers for the following species: Neisseria sicca, Neisseria polysaccharea and Neisseria meningitidis. There was a perfect correlation between the tetM genotype and the tetracycline susceptibility pattern of the Neisseria gonorrhoeae strains tested. The internal probe specific to N. gonorrhoeae SEQ ID NO. 2166 described in Example 31 can discriminate Neisseria gonorrhoeae from the other Neisseria sp.

The sensitivity of the assay with 40-cycle PCR was verified with two tetracycline resistant strains of N. gonorrhoeae. The detection limit was 5 copies of genomic DNA for both strains.

EXAMPLE 39:

Development of a PCR assay for the detection and identification of Shigella sp. and their associated trimethoprim resistance gene dhfrla. The analysis of publicly available dhfrla and other dhfr sequences revealed regions allowing the design of PCR primers specific to dhfrla sequences. The PCR primer pair (SEQ ID NOs. 1459 and 1460) was used in multiplex with the Escherichia coli/Shigella sp.specific primers SEQ ID NOs. 1661 and 1665 described in Example 28. Sequence alignment analysis of dhfrla sequences revealed regions suitable for the design of an internal probe specific to dhfrIa (SEQ ID NO. 2253). PCR amplification and agarose gel analysis of the amplified products were performed as described in Example 28 with an annealing temperature of 60 °C. The specificity of the multiplex assay with 40-cycle PCR was verified by using 0.1 ng of purified genomic DNA from a panel of bacteria. The list included the following trimethoprim-sensitive strains, Salmonella typhimyurium, Salmonella typhi, Salmonella enteritidis, Tatumella ptyseos, Klebsiella pneumoniae, Enterobacter aerogenes, Citrobacter farmeri, Campylobacter jejuni, Serratia marcescens, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, six trimethoprim-resistant Escherichia coli strains (containing dhfrla or dhfrVII or dhfrVII or dhfrXII or

dhfrXIII or dhfrXV), four trimethoprim-resistant strains containing dhfrIa (Shigella sonnei, Shigella flexneri, Shigella dysenteriae and Escherichia coli). There was a perfect correlation between the dhfrIa genotype and the trimethoprim susceptibility pattern of the Escherichia coli and Shigella sp. strains tested. The dhfrIa primers were specific to the dhfrIa gene and did not amplify any of the other trimethoprim-resistant dhfr genes tested. The sensitivity of the multiplex assay with 40-cycle PCR was verified with three strains of trimethoprim-resistant strains of Shigella sp. The detection limit was 5 to 10 genome copies of DNA, depending on the Shigella sp. strains tested.

EXAMPLE 40:

Development of a PCR assay for the detection and identification of Acinetobacter baumannii and its associated aminoglycoside resistance gene aph(3')-VIa. The comparison of publicly available aph(3')-VIa sequence revealed regions allowing the design of PCR primers specific to aph(3')-VIa. The PCR primer pair (SEQ ID NOs. 1404 and 1405) was used in multiplex with the Acinetobacter baumanniispecific primers SEQ ID NOs. 1692 and 1693 described in Example 30. Analysis of the aph(3')-Vla sequence revealed region suitable for the design of an internal probe specific to aph(3')-VIa (SEQ ID NO. 2252). PCR amplification and agarose gel analysis of the amplified products were performed as described in Example 28. The specificity of the multiplex assay with 40-cycle PCR was verified by using 0.1 ng of purified genomic DNA from a panel of bacteria including: two aminoglycoside-resistant A. baumanni strains (containing aph(3')-VIa), one aminoglycoside-sensitive A. baumani strain, one of each of the following aminoglycoside-resistant bacteria, one Serratia marcescens strain containing the aminoglycoside-resistant gene aacC1, one Serratia marcescens strain containing the aminoglycoside-resistant gene aacC4, one Enterobacter cloacae strain containing the aminoglycoside-resistant gene aacC2, one Enterococcus faecalis containing the aminoglycoside-resistant gene aacA-aphD, one Pseudomonas

aeruginosa strain containing the aminoglycoside-resistant gene aac6IIa and one of each of the following aminoglycoside-sensitive bacterial species, Acinetobacter anitratus, Acinetobacter lwoffi, Psychobbacter phenylpyruvian, Neisseria gonorrhoeae, Haemophilus haemolyticus, Haemophilus influenzae, Yersinia enterolitica, Proteus vulgaris, Eikenella corrodens, Escherichia coli. There was a perfect correlation between the aph(3')-VIa genotype and the aminoglycoside susuceptibility pattern of the A. baumannii strains tested. The aph(3')-VIa-specific primers were specific to the aph(3')-VIa gene and did not amplify any of the other aminoglycoside-resistant genes tested. The sensitivity of the multiplex assay with 40-cycle PCR was verified with two strains of aminoglycoside-resistant strains of A. baumannii. The detection limit was 5 genome copies of DNA for both A. baumannii strains tested.

EXAMPLE 41:

Specific identification of Bacteroides fragilis using atpD (V-type) sequences. The comparison of atpD (V-type) sequences from a variety of bacterial species allowed the selection of PCR primers for Bacteroides fragilis. The strategy used to design the PCR primers was based on the analysis of a multiple sequence alignement of various atpD sequences from B. fragilis, as well as atpD sequences from the related species B. dispar, bacterial genera and archaea, especially representatives with phylogenetically related atpD sequences. A careful analysis of this alignment allowed the selection of oligonucleotide sequences which are conserved within the target species but which discriminate sequences from other species, especially from closely related species B. dispar, thereby permitting the species-specific and ubiquitous detection and identification of the target bacterial species.

The chosen primer pair, SEQ ID NOs. 2134-2135, produces an amplification product of 231 bp. Standard PCR was carried out on PTC-200 thermocyclers (MJ Research Inc.) using $0.4\mu M$ of each primers pair as described in Example 28. The

optimal cycling conditions for maximum sensitivity and specificity were as follows: three minutes at 95°C for initial denaturation, then forty cycles of two steps consisting of 1 second at 95°C and 30 seconds at 60°C, followed by terminal extension at 72°C for 2 minutes.

The format of this assay is not limited to the one described above. A person skilled in the art could adapt the assay for different formats such as PCR with real-time detection using molecular beacon probes. Molecular beacon probes designed to be used in this assay include, but are not limited to, SEQ ID NO. 2136 for the detection of the *B. fragilis* amplicon.

EXAMPLE 42:

Evidence for horizontal gene transfer in the evolution of the elongation factor Tu in Enterococci.

ABSTRACT

The elongation factor Tu, encoded by tuf genes, is a GTP binding protein that plays a central role in protein synthesis. One to three tuf genes per genome are present depending on the bacterial species. Most low G+C gram-positive bacteria carry only one tuf gene. We have designed degenerate PCR primers derived from consensus sequences of the tuf gene to amplify partial tuf sequences from 17 enterococcal species and other phylogenetically related species. The amplified DNA fragments were sequenced either by direct sequencing or by sequencing cloned inserts containing putative amplicons. Two different tuf genes (tufA and tufB) were found in 11 enterococcal species, including Enterococcus avium, E. casseliflavus, E. dispar, E. durans, E. faecium, E. gallinarum, E. hirae, E. malodoratus, E. mundtii, E. pseudoavium, and E. raffinosus. For the other six enterococcal species (E. cecorum, E. columbae, E. faecalis, E. sulfureus, E.

rRNA gene sequence analysis, the 11 species having two tuf genes all share a common ancestor, while the six species having only one copy diverged from the enterococcal lineage before that common ancestor. The presence of one or two copies of the tuf gene in enterococci was confirmed by Southern hybridization. Phylogenetic analysis of tuf sequences demonstrated that the enterococcal tufA gene branches with the Bacillus, Listeria and Staphylococcus genera, while the enterococcal tufB gene clusters with the genera Streptococcus and Lactococcus. Primary structure analysis showed that four amino acid residues within the sequenced regions are conserved and unique to the enterococcal tufB genes and the tuf genes of streptococci and L. lactis. The data suggest that an ancestral streptococcus or a streptococcus-related species may have horizontally transferred a tuf gene to the common ancestor of the 11 enterococcal species which now carry two tuf genes.

INTRODUCTION

The elongation factor Tu (EF-Tu) is a GTP binding protein playing a central role in protein synthesis. It mediates the recognition and transport of aminoacyl-tRNAs and their positioning to the A-site of the ribosome. The highly conserved function and ubiquitous distribution render the elongation factor a valuable phylogenetic marker among eubacteria and even throughout the archaebacterial and eukaryotic kingdoms. The *tuf* genes encoding elongation factor Tu are present in various copy numbers per bacterial genome. Most gram-negative bacteria contain two *tuf* genes. As found in *Escherichia coli*, the two genes, while being almost identical in sequence, are located in different parts of the bacterial chromosome. However, recently completed microbial genomes revealed that only one *tuf* gene is found in *Helicobacter pylori* as well as in some obligate parasitic bacteria, such as *Borrelia burgdorferi*, *Rickettsia prowazekii*, and *Treponema pallidum*, and in some cyanobacteria. In most gram-positive bacteria studied so far, only one *tuf* gene was found. However, Southern hybridization showed that there are two *tuf* genes in

some clostridia as well as in *Streptomyces coelicolor* and *S. lividans*. Up to three tuf-like genes have been identified in *S. ramocissimus*.

Although massive prokaryotic gene transfer is suggested to be one of the factors responsible for the evolution of bacterial genomes, the genes encoding components of the translation machinery are thought to be highly conserved and difficult to be transferred horizontally due to the complexity of their interactions. However, a few recent studies demonstrated evidence that horizontal gene transfer has also occurred in the evolution of some genes coding for the translation apparatus, namely, 16S rRNA and some aminoacyl-tRNA synthetases. No further data suggest that such a mechanism is involved in the evolution of the elongation factors. Previous studies concluded that the two copies of *tuf* genes in the genomes of some bacteria resulted from an ancient event of gene duplication. Moreover, a study of the *tuf* gene in *R. prowazekii* suggested that intrachromosomal recombination has taken place in the evolution of the genome of this organism.

To date, little is known about the *tuf* genes of enterococcal species. In this study, we analyzed partial sequences of *tuf* genes in 17 enterococcal species, namely, *E. avium*, *E. casseliflavus*, *E. cecorum*, *E. columbae*, *E. dispar*, *E. durans*, *E. faecalis*, *E. faecium*, *E. gallinarum*, *E. hirae*, *E. malodoratus*, *E. mundtii*, *E. pseudoavium*, *E. raffinosus*, *E. saccharolyticus*, *E. solitarius*, and *E. sulfureus*. We report here the presence of two divergent copies of *tuf* genes in 11 of these enterococcal species. The 6 other species carried a single *tuf* gene. The evolutionary implications are discussed.

MATERIALS AND METHODS

Bacterial strains. Seventeen enterococcal strains and other gram-positive bacterial strains obtained from the American Type Culture Collection (ATCC, Manassas, Va.) were used in this study (Table 16). All strains were grown on sheep blood agar or in brain-heart infusion broth prior to DNA isolation.

DNA isolation. Bacterial DNAs were prepared using the G NOME DNA extraction kit (Bio101, Vista, Calif.) as previously described.

Sequencing of putative tuf genes. In order to obtain the tuf gene sequences of enterococci and other gram-positive bacteria, two sequencing approaches were used: 1) sequencing of cloned PCR products and 2) direct sequencing of PCR products. A pair of degenerate primers (SEQ ID NOs. 664 and 697) were used to amplify an 886-bp portion of the tuf genes from enterococcal species and other gram-positive bacteria as previously described. For E. avium, E. casseliflavus, E. dispar, E. durans, E. faecium, E. gallinarum, E. hirae, E. mundtii, E. pseudoavium, and E. raffinosus, the amplicons were cloned using the Original TA cloning kit (Invitrogen, Carlsbad, Calif.) as previously described. Five clones for each species were selected for sequencing. For E. cecorum, E. faecalis, E. saccharolyticus, and E. solitarius as well as the other gram-positive bacteria, the sequences of the 886bp amplicons were obtained by direct sequencing. Based on the results obtained from the earlier rounds of sequencing, two pairs of primers were designed for obtaining the partial tuf sequences from the other enterococcal species by direct sequencing. One pair of primers (SEQ ID NOs. 543 and 660) were used to amplify the enterococcal tuf gene fragments from E. columbae, E. malodoratus, and E. sulfureus. Another pair of primers (SEQ ID NOs. 664 and 661) were used to amplify the second tuf gene fragments from E. avium, E. malodoratus, and E. pseudoavium.

Prior to direct sequencing, PCR products were electrophoresed on 1% agarose gel at 120V for 2 hours. The gel was then stained with 0.02% methylene blue for 30 minutes and washed twice with autoclaved distilled water for 15 minutes. The gel slices containing PCR products of the expected sizes were cut out and purified with the QIAquick gel extraction kit (QIAgen Inc., Mississauga, Ontario, Canada) according to the manufacturer's instructions. PCR mixtures for sequencing were prepared as described previously. DNA sequencing was carried out with the Big DyeTM Terminator Ready Reaction cycle sequencing kit using a 377 DNA sequencer (PE Applied Biosystems, Foster City, Calif.). Both strands of the

amplified DNA were sequenced. The sequence data were verified using the SequencerTM 3.0 software (Gene Codes Corp., Ann Arbor, Mich.).

Sequence analysis and phylogenetic study. Nucleotide sequences of the tuf genes and their respective flanking regions for E. faecalis, Staphylococcus aureus, and Streptococcus pneumoniae, were retrieved from the TIGR microbial genome database and S. pyogenes from the University of Oklahoma database. DNA sequences and deduced protein sequences obtained in this study were compared with those in all publicly available databases using the BLAST and FASTA programs. Unless specified, sequence analysis was conducted with the programs from GCG package (Version 10; Genetics Computer Group, Madison, Wisc.). Sequence alignment of the tuf genes from 74 species representing all three kingdoms of life (Tables 16 and 17) were carried out by use of Pileup and corrected upon visual analysis. The N- and C-termini extremities of the sequences were trimmed to yield a common block of 201 amino acids sequences and equivocal residues were removed. Phylogenetic analysis was performed with the aid of PAUP 4.0b4 written by Dr. David L. Swofford (Sinauer Associates, Inc., Publishers, Sunderland, Mass.). The distance matrix and maximum parsimony were used to generate phylogenetic trees and bootstrap resampling procedures were performed using 500 and 100 replications in each analysis, respectively.

Protein structure analysis. The crystal structures of (i) Thermus aquaticus EF-Tu in complex with Phe-tRNA^{Phe} and a GTP analog and (ii) E. coli EF-Tu in complex with GDP served as templates for constructing the equivalent models for enterococcal EF-Tu. Homology modeling of protein structure was performed using the SWISS-MODEL server and inspected using the SWISS-PDB viewer version 3.1.

Southern hybridization. In a previous study, we amplified and cloned an 803-bp PCR product of the *tuf* gene fragment from *E. faecium*. Two divergent sequences of the inserts, which we assumed to be *tufA* and *tufB* genes, were obtained. The recombinant plasmid carrying either *tufA* or *tufB* sequence was used to generate two probes labeled with Digoxigenin (DIG)-11-dUTP by PCR

incorporation following the instructions of the manufacturer (Boehringer Mannheim, Laval, Québec, Canada). Enterococcal genomic DNA samples (1-2 μg) were digested to completion with restriction endonucleases BglII and XbaI as recommended by the supplier (Amersham Pharmacia Biotech, Mississauga, Ontario, Canada). These restriction enzymes were chosen because no restriction sites were observed within the amplified tuf gene fragments of most enterococci. Southern blotting and filter hybridization were performed using positively charged nylon membranes (Boehringer Mannheim) and QuikHyb hybridization solution (Stratagene Cloning Systems, La Jolla, Calif.) according to the manufacturers' instructions with modifications. Twenty µl of each digestion were electrophoresed for 2 h at 120V on a 0.8% agarose gel. The DNA fragments were denatured with 0.5 M NaOH and transferred by Southern blotting onto a positively charged nylon membrane (Boehringer Mannheim). The filters were pre-hybridized for 15 min and then hybridized for 2 h in the QuikHyb solution at 68°C with either DIG-labeled probe. Posthybridization washings were performed twice with 0.5x SSC, 1% SDS at room temperature for 15 min and twice in the same solution at 60°C for 15 min. Detection of bound probes was achieved using disodium 3- (4-methoxyspiro (1,2dioxetane-3,2'- (5'-chloro) tricyclo(3,3.1.1^{3.7}) decan)-4-yl) phenyl phosphate (CSPD) (Boehringer Mannheim) as specified by the manufacturer.

GenBank submission. The GenBank accession numbers for partial tuf gene sequences generated in this study are given in Table 16.

RESULTS

Sequencing and nucleotide sequence analysis. In this study, all gram-positive bacteria other than enterococci yielded a single tuf sequence of 886 bp using primers SEQ ID NOs. 664 and 697 (Table 16). Each of four enterococcal species including E. cecorum, E. faecalis, E. saccharolyticus, and E. solitarius also yielded one 886-bp tuf sequence. On the other hand, for E. avium, E. casseliflavus, E. dispar, E. durans, E. faecium, E. gallinarum, E. hirae, E. mundtii, E. pseudoavium,

and E. raffinosus, direct sequencing of the 886-bp fragments revealed overlapping peaks according to their sequence chromatograms, suggesting the presence of additional copies of the tuf gene. Therefore, the tuf gene fragments of these 10 species were cloned first and then sequenced. Sequencing data revealed that two different types of tuf sequences (tufA and tufB) are found in eight of these species including E. casseliflavus, E. dispar, E. durans, E. faecium, E. gallinarum, E. hirae, E. mundtii, and E. raffinosus. Five clones from E. avium and E. pseudoavium yielded only a single tuf sequence. These new sequence data allowed the design of new primers specific for the enterococcal tufA or tufB sequences. Primers SEQ ID NOs. 543 and 660 were designed to amplify only enterococcal tufA sequences and a 694-bp fragment was amplified from all 17 enterococcal species. The 694-bp sequences of tufA genes from E. columbae, E. malodoratus, and E. sulfureus were obtained by direct sequencing using these primers. Primers SEO ID NOs. 664 and 661 were designed for the amplification of 730-bp portion of tufB genes and yielded the expected fragments from 11 enterococcal species, including E. malodoratus and the 10 enterococcal species in which heterogeneous tuf sequences were initially found. The sequences of the tufB fragments for E. avium, E. malodoratus and E. pseudoavium were determined by direct sequencing using the primers SEQ ID NOs. 664 and 661. Overall, tufA gene fragments were obtained from all 17 enterococcal species but tufB gene fragments were obtained with only 11 enterococcal species (Table 16).

The identities between tufA and tufB for each enterococcal species were 68-79% at the nucleotide level and 81 to 89% at the amino acid level. The tufA gene is highly conserved among all enterococcal species with identities varying from 87% to 99% for DNA and 93% to 99% for amino acid sequences, while the identities among tufB genes of enterococci varies from 77% to 92% for DNA and 91% to 99% for amino acid sequences, indicating their different origins and evolution (Table 18). Since E. solitarius has been transferred to the genus Tetragenococcus, which is also a low G+C gram-positive bacterium, our sequence comparison did not include this species as an enterococcus. G+C content of enterococcal tufA

sequences ranged from 40.8% to 43.1%, while that of enterococcal tufB sequences varied from 37.8% to 46.3%. Based on amino acid sequence comparison, the enterococcal tufA gene products share higher identities with those of Abiotrophia adiacens, Bacillus subtilis, Listeria monocytogenes, S. aureus, and S. epidermidis. On the other hand, the enterococcal tufB gene products share higher percentages of amino acid identity with the tuf genes of S. pneumoniae, S. pyogenes and Lactococcus lactis (Table 18).

In order to elucidate whether the two enterococcal tuf sequences encode genuine EF-Tu, the deduced amino acid sequences of both genes were aligned with other EF-Tu sequences available in SWISSPROT (Release 38). Sequence alignment demonstrated that both gene products are highly conserved and carry all conserved residues present in this portion of prokaryotic EF-Tu (Figure 4). Therefore, it appears that both gene products could fulfill the function of EF-Tu. The partial tuf gene sequences encode the portion of EF-Tu from residues 117 to 317, numbered as in $E.\ coli$. This portion makes up of the last four α -helices and two β -strands of domain I, the entire domain II and the N-terminal part of domain III on the basis of the determined structures of $E.\ coli$ EF-Tu.

Based on the deduced amino acid sequences, the enterococcal *tufB* genes have unique conserved residues Lys129, Leu140, Ser230, and Asp234 (*E. coli* numbering) that are also conserved in streptococci and *L. lactis*, but not in the other bacteria (Figure 4). All these residues are located in loops except for Ser230. In other bacteria the residue Ser230 is substituted for highly conserved Thr, which is the 5th residue of the third β-strand of domain II. This region is partially responsible for the interaction between the EF-Tu and aminoacyl-tRNA by the formation of a deep pocket for any of the 20 naturally occurring amino acids. According to our three-dimensional model (data not illustrated), the substitution Thr230→Ser in domain II of EF-Tu may have little impact on the capability of the pocket to accommodate any amino acid. However, the high conservation of Thr230 comparing to the unique Ser substitution found only in streptococci and 11 enterococci could suggest a subtle functional role for this residue.

The tuf gene sequences obtained for E. faecalis, S. aureus, S. pneumoniae and S. pyogenes were compared with their respective incomplete genome sequence. Contigs with more than 99% identity were identified. Analysis of the E. faecalis genome data revealed that the single E. faecalis tuf gene is located within an str operon where tuf is preceded by fus that encodes the elongation factor G. This str operon is present in S. aureus and B. subtilis but not in the two streptococcal genomes examined. The 700-bp or so sequence upstream the S. pneumoniae tuf gene has no homology with any known gene sequences. In S. pyogenes, the gene upstream of tuf is similar to a cell division gene, ftsW, suggesting that the tuf genes in streptococci are not arranged in a str operon.

Phylogenetic analysis. Phylogenetic analysis of the *tuf* amino acid sequences with representatives of eubacteria, archeabacteria, and eukaryotes using neighborjoining and maximum parsimony methods showed three major clusters representing the three kingdoms of life. Both methods gave similar topologies consistent with the rRNA gene data (data not shown). Within the bacterial clade, the tree is polyphyletic but *tufA* genes from all enterococcal species always clustered with those from other low G+C gram-positive bacteria (except for streptococci and lactococci), while the *tufB* genes of the 11 enterococcal species form a distinct cluster with streptococci and *L. lactis* (Figure 5). Duplicated genes from the same organism do not cluster together, thereby not suggesting evolution by recent gene duplication.

Southern hybridization. Southern hybridization of BglII/XbaI digested genomic DNA from 12 enterococcal species tested with the tufA probe (DIG-labeled tufA fragment from E. faecium) yielded two bands of different sizes in 9 species, which also carried two divergent tuf sequences according to their sequencing data. For E. faecalis and E. solitarius, a single band was observed indicating that one tuf gene is present (Figure 6). A single band was also found when digested genomic DNA from S. aureus, S. pneumoniae, and S. pyogenes were hybridized with the tufA probe (data not shown). For E. faecium, the presence of three bands can be explained by the existence of a XbaI restriction site in the

middle of the *tufA* sequence, which was confirmed by sequencing data. Hybridization with the *tufB* probe (DIG-labeled *tufB* fragment of *E. faecium*) showed a banding profile similar to the one obtained with the *tufA* probe (data not shown).

DISCUSSION

In this study, we have shown that two divergent copies of genes encoding the elongation factor Tu are present in some enterococcal species. Sequence data revealed that both genes are highly conserved at the amino acid level. One copy (tufA) is present in all enterococcal species, while the other (tufB) is present only in 11 of the 17 enterococcal species studied. Based on 16S rRNA sequence analysis, these 11 species are members of three different enterococcal subgroups (E. avium, E. faecium, and E. gallinarum species groups) and a distinct species (E. dispar). Moreover, 16S rDNA phylogeny suggests that these 11 species possessing 2 tuf genes all share a common ancestor before they further evolved to become the modern species. Since the six other species having only one copy diverged from the enterococcal lineage before that common ancestor, it appears that the presence of one tuf gene in these six species is not attributable to gene loss.

Two clusters of low G+C gram-positive bacteria were observed in the phylogenetic tree of the *tuf* genes: one contains a majority of low G+C gram-positive bacteria and the other contains lactococci and streptococci. This is similar to the finding on the basis of phylogenetic analysis of the 16S rRNA gene and the *hrcA* gene coding for a unique heat-shock regulatory protein. The enterococcal *tufA* genes branched with most of the low G+C gram-positive bacteria, suggesting that they originated from a common ancestor. On the other hand, the enterococcal *tufB* genes branched with the genera *Streptococcus* and *Lactococcus* that form a distinct lineage separated from other low G+C gram-positive bacteria (Figure 5). The finding that these EF-Tu proteins share some conserved amino acid residues unique to this branch also supports the idea that they may share a common ancestor. Although these conserved residues might result from convergent

even for a few residues, seems to be rare, making it an unlikely event. Moreover, no currently known selective pressure, if any, would account for keeping one versus two *tuf* genes in bacteria. The G+C contents of enterococcal *tufA* and *tufB* sequences are similar, indicating that they both originated from low G+C grampositive bacteria, in accordance with the phylogenetic analysis.

The tuf genes are present in various copy numbers in different bacteria. Furthermore, the two tuf genes are normally associated with characteristic flanking genes. The two tuf gene copies commonly encountered within gram-negative bacteria are part of the bacterial str operon and tRNA-tufB operon, respectively. The arrangement of tufA in the str operon was also found in a variety of bacteria, including Thermotoga maritima, the most ancient bacteria sequenced so far, Aquifex aeolicus, cyanobacteria, Bacillus sp., Micrococcus luteus, Mycobacterium tuberculosis, and Streptomyces sp. Furthermore, the tRNA-tufB operon has also been identified in Aquifex aeolicus, Thermus thermophilus, and Chlamydia trachomatis. The two widespread tuf gene arrangements argue in favor of their ancient origins. It is noteworthy that most obligate intracellular parasites, such as Mycoplasma sp., R. prowazekii, B. burgdorferi, and T. pallidum, contain only one tuf gene. Their flanking sequences are distinct from the two conserved patterns as a result of selection for effective propagation by an extensive reduction in genome size by intragenomic recombination and rearrangement.

Most gram-positive bacteria with low G+C content sequenced to date contain only a single copy of the tuf gene as a part of the str operon. This is the case for B. subtilis, S. aureus and E. faecalis. PCR amplification using a primer targeting a conserved region of the fus gene and the tufA-specific primer SEQ ID NO. 660, but not the tufB-specific primer SEQ ID NO. 661, yielded the expected amplicons for all 17 enterococcal species tested, indicating the presence of the fus-tuf organization in all enterococci (data not shown). However, in the genomes of S. pneumoniae and S. pyogenes, the sequences flanking the tuf genes varies although the tuf gene itself remains highly conserved. The enterococcal tufB genes are

clustered with streptococci, but at present we do not have enough data to identify the genes flanking the enterococcal *tufB* genes. Furthermore, the functional role of the enterococcal *tufB* genes remains unknown. One can only postulate that the two divergent gene copies are expressed under different conditions.

The amino acid sequence identities between the enterococcal tufA and tufB genes are lower than either i) those between the enterococcal tufA and the tuf genes from other low G+C gram-positive bacteria (streptococci and lactococci excluded) or ii) those between the enterococcal tufB and streptococcal and lactococcal tuf genes. These findings suggest that the enterococcal tufA genes share a common ancestor with other low G+C gram-positive bacteria via the simple scheme of vertical evolution, while the enterococcal tufB genes are more closely related to those of streptococci and lactococci. The facts that some enterococci possess an additional tuf gene and that the single streptococcal tuf gene is not clustered with other low G+C gram-positive bacteria cannot be explained by the mechanism of gene duplication or intrachromosomal recombination. According to sequence and phylogenetic analysis, we propose that the presence of the additional copy of the tuf genes in 11 enterococcal species is due to horizontal gene transfer. The common ancestor of the 11 enterococcal species now carrying tufB genes acquired a tuf gene from an ancestral streptococcus or a streptococcus-related species during enterococcal evolution through gene transfer before the diversification of modern enterococci. Further study of the flanking regions of the gene may provide more clues for the origin and function of this gene in enterococci.

Recent studies of genes and genomes have demonstrated that considerable horizontal transfer occurred in the evolution of aminoacyl-tRNA synthetases in all three kingdoms of life. The heterogeneity of 16S rRNA is also attributable to horizontal gene transfer in some bacteria, such as *Streptomyces*, *Thermomonospora chromogena* and *Mycobacterium celatum*. In this study, we provide the first example in support of a likely horizontal transfer of the *tuf* gene encoding the elongation factor Tu. This may be an exception since stringent functional constraints do not allow for frequent horizontal transfer of the *tuf* gene as with

other genes. However, enterococcal tuf genes should not be the only such exception as we have noticed that the phylogeny of Streptomyces tuf genes is equally or more complex than that of enterococci. For example, the three tuf-like genes in a high G+C gram-positive bacterium, S. ramocissimus, branched with the tuf genes of phylogenetically divergent groups of bacteria (Figure 5). Another example may be the tuf genes in clostridia, which represent a phylogenetically very broad range of organisms and form a plethora of lines and groups of various complexities and depths. Four species belonging to three different clusters within the genus Clostridium have been shown by Southern hybridization to carry two copies of the tuf gene. Further sequence data and phylogenetic analysis may help interpreting the evolution of the elongation factor Tu in these gram-positive bacteria. Since the tuf genes and 16S rRNA genes are often used for phylogenetic study, the existence of duplicate genes originating from horizontal gene transfer may alter the phylogeny of microorganisms when the laterally acquired copy of the gene is used for such analysis. Hence, caution should be taken in interpreting phylogenetic data. In addition, the two tuf genes in enterococci have evolved separately and are distantly related to each other phylogenetically. The enterococcal tufB genes are less conserved and unique to the 11 enterococcal species only. We previously demonstrated that the enterococcal tufA genes could serve as a target to develop a DNA-based assay for identification of enterococci. The enterococcal tufB genes would also be useful in identification of these 11 enterococcal species.

EXAMPLE 43:

Elongation Factor Tu (tuf) and the F-ATPase beta-subunit (atpD) as phylogenetic tools for species of the family Enterobacteriaceae.

SUMMARY

The phylogeny of enterobacterial species commonly found in clinical samples was analyzed by comparing partial sequences of their elongation factor Tu (tuf) genes and their F-ATPase beta-subunit (atpD) genes. A 884-bp fragment for tuf and a 884- or 871-bp fragment for atpD were sequenced for 88 strains of 72 species from 25 enterobacterial genera. The atpD sequence analysis revealed a specific indel to Pantoea and Tatumella species showing for the first time a tight phylogenetic affiliation between these two genera. Comprehensive tuf and atpD phylogenetic trees were constructed and are in agreement with each other. Monophyletic genera are Yersinia, Pantoea, Edwardsiella, Cedecea, Salmonella, Serratia, Proteus, and Providencia. Analogous trees were obtained based on available 16S rDNA sequences from databases. tuf and atpD phylogenies are in agreement with the 16S rDNA analysis despite the smaller resolution power for the latter. In fact, distance comparisons revealed that tuf and atpD genes provide a better resolution for pairs of species belonging to the family Enterobacteriaceae. However, 16S rDNA distances are better resolved for pairs of species belonging to different families. In conclusion, tuf and atpD conserved genes are sufficiently divergent to discriminate different species inside the family Enterobacteriaceae and offer potential for the development of diagnostic tests based on DNA to identify enterobacterial species.

INTRODUCTION

Members of the family *Enterobacteriaceae* are facultatively anaerobic gramnegative rods, catalase-positive and oxydase-positive (Brenner, 1984). They are found in soil, water, plants, and in animals from insects to man. Many enterobacteria are opportunistic pathogens. In fact, members of this family are responsible for about 50 % of nosocomial infections in the United States (Brenner, 1984). Therefore, this family is of considerable clinical importance.

Major classification studies on the family Enterobacteriaceae are based on phenotypic traits (Brenner et al., 1999; Brenner et al., 1980; Dickey & Zumoff,

1988; Farmer III et al., 1980; Farmer III et al., 1985b; Farmer III et al., 1985a) such as biochemical reactions and physiological characteristics. However, phenotypically distinct strains may be closely related by genotypic criteria and may belong to the same genospecies (Bercovier et al., 1980; Hartl & Dykhuizen, 1984). Also, phenotypically close strains (biogroups) may belong to different genospecies, like Klebsiella pneumoniae and Enterobacter aerogenes (Brenner, 1984) for example. Consequently, identification and classification of certain species may be ambiguous with techniques based on phenotypic tests (Janda et al., 1999; Kitch et al., 1994; Sharma et al., 1990).

More advances in the classification of members of the family Enterobacteriaceae have come from DNA-DNA hybridization studies (Brenner et al., 1993; Brenner et al., 1986; Brenner, et al., 1980; Farmer III, et al., 1980; Farmer III, et al., 1985b; Izard et al., 1981; Steigerwalt et al., 1976). Furthermore, the phylogenetic significance of bacterial classification based on 16S rDNA sequences has been recognized by many workers (Stackebrandt & Goebel, 1994; Wayne et al., 1987). However, members of the family Enterobacteriaceae have not been subjected to extensive phylogenetic analysis of 16S rDNA (Sproer et al., 1999). In fact, this molecule was not thought to solve taxonomic problems concerning closely related species because of its very high degree of conservation (Brenner, 1992; Sproer, et al., 1999). Another drawback of the 16S rDNA gene is that it is found in several copies within the genome (seven in Escherichia coli and Salmonella typhimurium) (Hill & Harnish, 1981). Due to sequence divergence between the gene copies, direct sequencing of PCR products is often not suitable to achieve a representative sequence (Cilia et al., 1996; Hill & Harnish, 1981). Other genes such as gap and ompA (Lawrence et al., 1991), rpoB (Mollet et al., 1997), and infB (Hedegaard et al., 1999) were used to resolve the phylogeny of enterobacteria. However, none of these studies covered an extensive number of species.

tuf and atpD are the genes encoding the elongation factor Tu (EF-Tu) and the F-ATPase beta-subunit, respectively. EF-Tu is involved in peptide chain formation (Ludwig et al., 1990). The two copies of the tuf gene (tufA and tufB) found in enterobacteria (Sela et al., 1989) share high identity level (99 %) in Salmonella typhimurium and in E. coli. The recombination phenomenon could explain sequence homogenization between the two copies (Abdulkarim & Hughes, 1996; Grunberg-Manago, 1996). F-ATPase is present on the plasma membranes of eubacteria (Nelson & Taiz, 1989). It functions mainly in ATP synthesis (Nelson & Taiz, 1989) and the beta-subunit contains the catalytic site of the enzyme. EF-Tu and F-ATPase are highly conserved throughout evolution and shows functional constancy (Amann et al., 1988; Ludwig, et al., 1990). Recently, phylogenies based on protein sequences from EF-Tu and F-ATPase beta-subunit showed good agreement with each other and with the rDNA data (Ludwig et al., 1993).

We elected to sequence 884-bp fragments of *tuf* and *atpD* from 88 clinically relevant enterobacterial strains representing 72 species from 25 genera. These sequences were used to create phylogenetic trees that were compared with 16S rDNA trees. These trees revealed good agreement with each others and demonstrated the high resolution of *tuf* and *atpD* phylogenies at the species level.

MATERIALS AND METHODS

Bacterial strains and genomic material. All bacterial strains used in this study were obtained from the American Type Culture Collection (ATCC) or the Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH (DSMZ). These enterobacteria can all be recovered from clinical specimens, but not all are pathogens. Whenever possible, we choose type strains. Identification of all strains was confirmed by classical biochemical tests using the automated system MicroScan WalkAway-96 system equipped with a Negative BP Combo Panel Type 15 (Dade Behring Canada). Genomic DNA was purified using the G NOME

DNA kit (Bio 101). Genomic DNA from Yersinia pestis was kindly provided by Dr. Robert R. Brubaker. Strains used in this study and their descriptions are shown in Table 19.

PCR primers. The eubacterial tuf and atpD gene sequences available from public databases were analyzed using the GCG package (version 8.0) (Genetics Computer Group). Based on multiple sequence alignments, two highly conserved regions were chosen for each genes, and PCR primers were derived from these regions with the help of Oligo primer analysis software (version 5.0) (National Biosciences). A second 5' primer was design to amplify the gene atpD for few enterobacteria difficult to amplify with the first primer set. When required, the primers contained inosines or degeneracies to account for variable positions. Oligonucleotide primers were synthesized with a model 394 DNA/RNA synthesizer (PE Applied Biosystems). PCR primers used in this study are listed in Table 20.

DNA sequencing. An 884-bp portion of the *tuf* gene and an 884-bp portion (or alternatively an 871-bp portion for a few enterobacterial strains) of the *atpD* gene were sequenced for all enterobacteria listed in the first strain column of Table 19. Amplification was performed with 4 ng of genomic DNA. The 40-μl PCR mixtures used to generate PCR products for sequencing contained 1·0 μM each primer, 200 μM each deoxyribonucleoside triphosphate (Pharmacia Biotech), 10 mM Tris-HCl (pH 9·0 at 25 °C), 50 mM KCl, 0·1 % (w/v) Triton X-100, 2·5 mM MgCl₂, 0·05 mM BSA, 0·3 U of *Taq* DNA polymerase (Promega) coupled with TaqStartTM antibody (Clontech Laboratories). The TaqStartTM neutralizing monoclonal antibody for *Taq* DNA polymerase was added to all PCR mixtures to enhance efficiency of amplification (Kellogg *et al.*, 1994). The PCR mixtures were subjected to thermal cycling (3 min at 95 °C and then 35 cycles of 1 min at 95 °C, 1 min at 55 °C for *tuf* or 50 °C for *atpD*, and 1 min at 72 °C, with a 7-min final extension at 72 °C) using a PTC-200 DNA Engine thermocycler (MJ Research).

PCR products having the predicted sizes were recovered from an agarose gel stained for 15 min with 0.02 % of methylene blue followed by washing in sterile distilled water for 15 min twice (Flores et al., 1992). Subsequently, PCR products having the predicted sizes were recovered from gels using the QIAquick gel extraction kit (QIAGEN).

Both strands of the purified amplicons were sequenced using the ABI Prism BigDye Terminator Cycle Sequencing Ready Reaction Kit (PE Applied Biosystems) on an automated DNA sequencer (Model 377). Amplicons from two independant PCR amplifications were sequenced for each strain to ensure the absence of sequencing errors attributable to nucleotide miscorporations by the *Taq* DNA polymerase. Sequence assembly was performed with the aid of Sequencher 3.0 software (Gene Codes).

Phylogenetic analysis. Multiple sequence alignments were performed using PileUp from the GCG package (Version 10.0) (Genetics Computer Group) and checked by eye with the editor SeqLab to edit sequences if necessary and to note which regions were to be excluded for phylogenetic analysis. Vibrio cholerae and Shewanella putrefaciens were used as outgroups. Bootstrap subsets (750 sets) and phylogenetic trees were generated with the Neighbor Joining algorithm from Dr. David Swofford's PAUP (Phylogenetic Analysis Using Parsimony) Software version 4.0b4 (Sinauer Associates) and with tree-bisection branch-swapping. The distance model used was Kimura (1980) two-parameter. Relative rate test was performed with the aid of Phyltest program version 2.0 (c).

RESULTS AND DISCUSSION

DNA amplification, sequencing and sequence alignments

A PCR product of the expected size of 884 bp was obtained for *tuf* and of 884 or 871 bp for *atpD* from all bacterial strains tested. After subtracting for biased

primer regions and ambiguous single strand data, sequences of at least 721 bp for tuf and 713 bp for atpD were submitted to phylogenetic analyses. These sequences were aligned with tuf and atpD sequences available in databases to verify that the nucleotide sequences indeed encoded a part of tested genes. Gaps were excluded to perform phylogenetic analysis.

Signature sequences

From the sequence alignments obtained from both tested genes, only one insertion was observed. This five amino acids insertion is located between the positions 325 and 326 of atpD gene of E. coli strain K-12 (Saraste et al., 1981) and can be considered a signature sequence of Tatumella ptyseos and Pantoea species (Fig. 7). The presence of a conserved indel of defined length and sequence and flanked by conserved regions could suggest a common ancestor, particularly when members of a given taxa share this indel (Gupta, 1998). To our knowledge, high relatedness between the genera Tatumella and Pantoea is demonstrated for the first time.

Enterobacter agglomerans ATCC 27989 sequence does not possess the five amino acid indel (Fig. 7). This indel could represent a useful marker to help resolve the Enterobacter agglomerans and Pantoea classification. Indeed, the transfer of Enterobacter agglomerans to Pantoea agglomerans was proposed in 1989 by Gavini et al. (Gavini et al., 1989). However, some strains are provisionally classified as Pantoea sp. until their interrelatedness is elucidated (Gavini, et al., 1989). Since the transfer was proposed, the change of nomenclature has not yet been made for all Enterobacter agglomerans in the ATCC database. The absence of the five amino acids indel suggests that some strains of Enterobacter agglomerans most likely do not belong to the genus Pantoea.

Phylogenetic trees based on partial tuf sequences, atpD sequences, and published 16S rDNA data of members of the Enterobacteriaceae.

Representative trees constructed from tuf and atpD sequences with the neighbor-joining method are shown in Fig. 8. The phylogenetic trees generated from partial tuf sequences and atpD sequences are very similar. Nevertheless, atpD tree shows more monophyletic groups corresponding to species that belong to the same genus. These groups are more consistent with the actual taxonomy. For both genes, some genera are not monophyletic. These results support previous phylogenies based on the genes gap and ompA (Lawrence, et al., 1991), rpoB (Mollet, et al., 1997), and infB (Hedegaard, et al., 1999) which all showed that the genera Escherichia and Klebsiella are polyphyletic. There were few differences in branching between tuf and atpD genes.

Even though Pantoea agglomerans and Pantoea dispersa indels were excluded for phylogenetic analysis, these two species grouped together and were distant from Enterobacter agglomerans ATCC 27989, adding another evidence that the latter species is heterogenous and that not all members of this species belong to the genus Pantoea. In fact, the E. agglomerans strain ATCC 27989 exhibits branch lengths similar to others Enterobacter species with both genes. Therefore, we suggest that this strain belong to the genus Enterobacter until further reclassification of that genus.

tuf and atpD trees exhibit very short genetic distances between taxa belonging to the same genetic species including species segregated for clinical considerations. This first concern E. coli and Shigella species that were confirmed to be the same genetic species by hybridization studies (Brenner et al., 1972; Brenner et al., 1972; Brenner et al., 1982) and phylogenies based on 16S rDNA (Wang et al., 1997) and rpoB genes (Mollet, et al., 1997). Hybridization studies (Bercovier, et al., 1980) and phylogeny based on 16S rDNA genes (Ibrahim et al., 1994) demonstrated also that Yersinia pestis and Y. pseudotuberculosis are the same genetic species. Among

Yersinia pestis and Y. pseudotuberculosis, the three Klebsiella pneumoniae subspecies, E. coli-Shigella species, and Salmonella choleraesuis subspecies, Salmonella is a less tightly knit species than the other genetic species. The same is true for E. coli and Shigella species.

Escherichia fergusonii is very close to E. coli-Shigella genetic species. This observation is corroborated by 16S rDNA phylogeny (McLaughlin et al., 2000) but not by DNA hybridization values. In fact, E. fergusonii is only 49% to 63% related to E. coli-Shigella (Farmer III, et al., 1985b). It was previously observed that very recently diverged species may not be recognizable based on 16S rDNA sequences although DNA hybridization established them as different species (Fox et al., 1992). Therefore, E. fergusonii could be a new "quasi-species".

atpD phylogeny revealed Salmonella subspecies divisions consistent with the actual taxonomy. This result was already observed by Christensen et al. (Christensen & Olsen, 1998). Nevertheless, tuf partial sequences discriminate less than atpD between Salmonella subspecies.

Overall, tuf and atpD phylogenies exhibit enough divergence between species to ensure efficient discrimination. Therefore, it could be easy to distinguish phenotypically close enterobacteria belonging to different genetic species such as Klebsiella pneumoniae and Enterobacter aerogenes.

Phylogenetic relationships between Salmonella, E. coli and C. freundii are not well defined. 16S rDNA and 23S rDNA sequence data reveals a closer relationship between Salmonella and E. coli than between Salmonella and C. freundii (Christensen et al., 1998), while DNA homology studies (Selander et al., 1996) and infB phylogeny (Hedegaard, et al., 1999) showed that Salmonella is more closely related to C. freundii than to E. coli. In that regard, tuf and atpD phylogenies are coherent with 16S rDNA and 23S rDNA sequence analysis.

Phylogenetic analyses were also performed using amino acids sequences. *tuf* tree based on amino acids is characterized by a better resolution between taxa outgroup and taxa ingroup (enterobacteria) than tree based on nucleic acids whereas *atpD* trees based on amino acids and nucleic acids give almost the same resolution between taxa outgroup and ingroup (data not shown).

Relative rate test (or two cluster test (Takezaki et al., 1995)) evaluates if evolution is constant between two taxa. Before to apply the test, the topology of a tree is determined by tree-building method without the assumption of rate constancy. Therefore, two taxa (or two groups of taxa) are compared with a third taxon that is an outgroup of the first two taxa (Takezaki, et al., 1995). Few pairs of taxa that exhibited a great difference between their branch lengths at particular nodes were chosen to perform the test. This test reveals that tuf and atpD are not constant in their evolution within the family Enterobacteriaceae. For tuf, for example, the hypothesis of rate constancy is rejected (Z value higher than 1.96) between Yersinia species. The same is true for Proteus species. For atpD, for example, evolution is not constant between Proteus species, between Proteus species and Providencia species, and between Yersinia species and Escherichia coli. For 16S rDNA, for example, evolution is not constant between two E. coli, between E. coli and Enterobacter aerogenes, and between E. coli and Proteus vulgaris. These results suggest that tuf, atpD and 16S rDNA could not serve as a molecular clock for the entire family Enterobacteriaceae.

Since the number and the nature of taxa can influence topology of trees, phylogenetic trees from tuf and atpD were reconstructed using sequences corresponding to strains for which 16S rDNA genes were published in GenEMBL. These trees were similar to those generated using 16S rDNA (Fig. 9). Nevertheless, 16S rDNA tree gave poorer resolution power than tuf and atpD gene trees. Indeed, these latter exhibited less multifurcation (polytomy) than the 16S rDNA tree.

Comparison of distances based on tuf, atpD, and 16S rDNA data.

tuf, atpD, and 16S rDNA distances (i.e. the number of differences per nucleotide site) were compared with each other for each pair of strains. We found that the tuf and atpD distances were respectively 2.268 ± 0.965 and 2.927 ± 0.896 times larger than 16S rDNA distances (Fig. 10a and b). atpD distances were 1.445 ± 0.570 times larger than tuf distances (Fig. 10c). Figure 10 also shows that the tuf, atpD, and 16S rDNA distances between members of different species of the same genus $(0.053 \pm 0.034, 0.060 \pm 0.020, \text{ and } 0.024 \pm 0.010, \text{ respectively})$ were in mean smaller than the distances between members of different genera belonging to the same family $(0.103 \pm 0.053, 0.129 \pm 0.051, \text{ and } 0.044 \pm 0.013, \text{ respectively}).$ However, the overlap exhibits with standard deviations add to a focus of evidences that some enterobacterial genera are not well defined (Brenner, 1984). In fact, many distances for pairs of species especially belonging to the genera Escherichia, Shigella, Enterobacter, Citrobacter, Klebsiella, and Kluyvera overlap distances for pairs of species belonging to the same genus (Fig. 10). For example, distances for pairs composed by species of Citrobacter and species of Klebsiella overlap distances for pairs composed by two Citrobacter or by two Klebsiella.

Observing the distance distributions, 16S rDNA distances reveal a clear separation between the families *Enterobacteriaceae* and *Vibrionaceae* despite the fact that the family *Vibrionaceae* is genetically very close to the *Enterobacteriaceae* (Fig. 10a and b). Nevertheless, *tuf* and *atpD* show higher discriminating power below the family level (Fig. 10a and b).

There were some discrepancies in the relative distances for the same pairs of taxa between the two genes studied. First, distances between Yersinia species are at least two times lower for atpD than for tuf (Fig. 10c). Also, distances at the family level (between Enterobacteriaceae and Vibrionaceae) show that Enterobacteriaceae is a tightlier knit family with atpD gene (Proteus genus

excepted) than with tuf gene. Both genes well delineate taxa belonging to the same species. There is one exception with atpD: Klebsiella planticola and K. ornithinolithica belong to the same genus but fit with taxa belonging to the same species (Fig. 10a and c). These two species are also very close genotypically with tuf gene. This suggest that Klebsiella planticola and K. ornithinolithica could be two newborn species. tuf and atpD genes exhibit little distances between Escherichia fergusonii and E. coli-Shigella species. Unfortunately, comparison with 16S rDNA could not be achieved because the E. fergusonii 16S rDNA sequence is not yet accessible in GenEMBL database. Therefore, the majority of phenotypically close enterobacteria could be easily discriminated genotypically using tuf and atpD gene sequences.

In conclusion, tuf and atpD genes exhibit phylogenies consistent with 16S rDNA genes phylogeny. For example, they reveal that the family Enterobacteriaceae is monophyletic. Moreover, tuf and atpD distances provide a higher discriminating power than 16S rDNA distances. In fact, tuf and atpD genes discriminate well between different genospecies and are conserved between strains of the same genetic species in such a way that primers and molecular probes for diagnostic purposes could be designed. Preliminary studies support these observations and diagnostic tests based on tuf and atpD sequence data to identify enterobacteria are currently under development.

EXAMPLE 44:

Testing new pairs of PCR primers selected from two species-specific genomic DNA fragments which are objects of our assigned US patent 6,001,564

Objective: The goal of these experiments is to demonstrate that it is relatively easy for a person skilled in the art to find other PCR primer pairs from the species-specific

fragments used as targets for detection and identification of a variety of microorganisms. In fact, we wish to prove that the PCR primers previously tested by our group and which are objects of the present patent application are not the only possible good choices for diagnostic purposes. For this example, we used diagnostic targets described in our assigned US patent 6,001,564.

Experimental strategy: We have selected randomly two species-specific genomic DNA fragments for this experiment. The first one is the 705-bp fragment specific to Staphylococcus epidermidis (SEQ ID NO: 36 from US patent 6,001,564) while the second one is the 466-bp fragment specific to Moraxella catarrhalis (SEQ ID NO: 29 from US patent 6,001,564). Subsequently, we have selected from these two fragments a number of PCR primer pairs other than those previously tested. We have chosen 5 new primer pairs from each of these two sequences which are well dispersed along the DNA fragment (Figures 11 and 12). We have tested these primers for their specificity and compared them with the original primers previously tested. For the specificity tests, we have tested all bacterial species closely related to the target species based on phylogenetic analysis with three conserved genes (rRNA genes, tuf and atpD). The rational for selecting a restricted number of bacterial species to evaluate the specificity of the new primer pairs is based on the fact that the lack of specificity of a DNA-based assay is attributable to the detection of closely related species which are more similar at the nucleotide level. Based on the phylogenetic analysis, we have selected (i) species from the closely related genus Staphylococcus, Enterococcus, Streptococcus and Listeria to test the specificity of the S. epidermidis-specific PCR assays and (ii) species from the closely related genus Moraxella, Kingella and Neisseria to test the specificity of the M. catarrhalisspecific PCR assays.

Materials and methods

Bacterial strains. All bacterial strains used for these experiments were obtained from the American Type Culture Collection (ATCC, Rockville, MD).

Genomic DNA isolation. Genomic DNA was purified from the ATCC reference strains by using the G-nome DNA kit (Bio 101 Inc., Vista, CA).

Oligonucleotide design and synthesis. PCR primers were designed with the help of the OligoTM primer analysis software Version 4.0 (National Biosciences Inc., Plymouth, Minn.) and synthesized using a model 391 DNA synthesizer (Applied Biosystems, Foster City, CA).

PCR assays. All PCR assays were performed by using genomic DNA purified from reference strains obtained from the ATCC. One μl of purified DNA preparation (containing 0.01 to 1 ng of DNA per μ l) was added directly into the PCR reaction mixture. The 20 µL PCR reactions contained final concentrations of 50 mM KCl, 10 mM Tris-HCl (pH 9.0), 0.1% Triton X-100, 2.5 mM MgCl₂, 0.4 μ M of each primer, 200 µM of each of the four dNTPs and 0.5 unit of Taq DNA polymerase (Promega, Madison, WI) combined with the TaqStartTM antibody (Clontech Laboratories Inc., Palo Alto, CA). An internal control was integrated into all amplification reactions to verify the efficiency of the amplification reaction as well as to ensure that significant PCR inhibition was absent. Primers amplifying a region of 252 bp from a control plasmid added to each amplification reaction were used to provide the internal control. PCR reactions were then subjected to thermal cycling (3 min at 95°C followed by 30 cycles of 1 second at 95°C for the denaturation step and 30 seconds at 50 to 65°C for the annealing-extension step) using a PTC-200 thermal cycler (MJ Research Inc., Watertown, MA). PCR amplification products were then analyzed by standard agarose gel (2%) electrophoresis. Amplification products were visualized in agarose gels containing 0.25 μ g/mL of ethidium bromide under UV at 254 nm.

Results

Tables 21 and 22 show the results of specificity tests with the 5 new primer pairs selected from SEQ ID NO: 29 (specific to *M. catarrhalis* from US patent 6,001,564) and SEQ ID NO: 36 (specific to *S. epidermidis* from US patent 6,001,564), respectively. In order to evaluate the performance of these new primers pairs, we compared them in parallel with the original primer pairs previously tested.

For *M. catarrhalis*, all of the 5 selected PCR primer pairs were specific for the target species because none of the closely related species could be amplified (Table 21). In fact, the comparison with the original primer pair SEQ ID NO: 118 + SEQ ID NO: 119 (from US patent 6,001,564) revaled that all new pairs showed identical results in terms of specificity and sensitivity thereby suggesting their suitability for diagnostic purposes.

For S. epidermidis, 4 of the 5 selected PCR primer pairs were specific for the target species (Table 22). It should be noted that for 3 of these four primer pairs the annealing temperature had to be increased from 55 °C to 60 or 65 °C to attain specificity for S. epidermidis. Again the comparison with the original primer pair SEQ ID NO: 145 + SEQ ID NO: 146 (from US patent 6,001,564) revealed that these four primer pairs were as good as the original pair. Increasing the annealing temperature for the PCR amplification is well known by persons skilled in the art to be a very effective way to improve the specificity of a PCR assay (Persing et al., 1993, Diagnostic Molecular Microbiology: Principles and Applications, American Society for Microbiology, Washington, D.C.; Ehrlich and Greenberg, 1994, PCRbased Diagnostics in Infectious Disease, Blackwell Scientific Publications, Boston, MA). In fact, those skilled in the art are well aware of the fact that the annealing temperature is critical for the optimization of PCR assays. Only the primer pair VBsep3 + VBsep4 amplified bacterial species other than S. epidermidis including the staphylococcal species S. capitis, S. cohnii, S. aureus, S. haemolyticus and S. hominis (Table 22). For this non-specific primer pair, increasing the annealing temperature

from 55 to 65 °C was not sufficient to attain the desired specificity. One possible explanation for the fact that it appears slightly easier to select species-specific primers for *M. catarrhalis* than for *S. epidermidis* is that *M. catarrhalis* is more isolated in phylogenetic trees than *S. epidermidis*. The large number of coagulase negative staphylococcal species such as *S. epidermidis* is largely responsible for this phylogenetic clustering.

Conclusion

These experiment clearly show that it is relatively easy for a person skilled in the art to select, from the species-specific DNA fragments selected as target for identification, PCR primer pairs suitable for diagnostic purposes other than those previously tested. The amplification conditions can be optimize by modifying critical variables such as the annealing temperature to attain the desired specificity and sensitivity. Consequently, we consider that it is legitimate to claim any possible primer sequences selected from the species-specific fragment and that it would be unfair to grant only the claims dealing with the primer pairs previously tested. By extrapolation, these results strongly suggest that it is also relatively easy for a person skilled in the art to select, from the species-specific DNA fragments, DNA probes suitable for diagnostic purposes other than those previously tested.

EXAMPLE 45:

Testing modified versions of PCR primers derived from the sequence of several primers which are objects of US patent 6,001,564.

Objective: The purpose of this project is to verify the efficiency of amplification by modified PCR primers derived from primers previously tested. The types of primer modifications to be tested include (i) variation of the sequence at one or more nucleotide positions and (ii) increasing or reducing the length of the primers. For this example, we used diagnostic targets described in US patent 6,001,564.

Experimental strategy:

a) Testing primers with nucleotide changes

We have designed 13 new primers which are derived from the *S. epidermidis*-specific SEQ ID NO: 146 from US patent 6,001,564 (Table 23). These primers have been modified at one or more nucleotide positions. As shown in Table 23, the nucleotide changes were introduced all along the primer sequence. Furthermore, instead of modifying the primer at any nucleotide position, the nucleotide changes were introduced at the third position of each codon to better reflect potential genetic variations *in vivo*. It should be noted that no nucleotide changes were introduced at the 3' end of the oligonucleotide primers because those skilled in the art are well aware of the fact that mimatches at the 3' end should be avoided (Persing *et al.*, 1993, Diagnostic Molecular Microbiology: Principles and Applications, American Society for Microbiology, Washington, D.C.). All of these modified primers were tested in PCR assays in combination with SEQ ID NO: 145 from US patent 6,001,564 and the efficiency of the amplification was compared with the original primer pair SEQ ID NO: 145 + SEQ ID NO: 146 previously tested in US patent 6,001,564.

b) Testing shorter or longer versions of primers

We have designed shorter and longer versions of the original S. epidermidis-specific PCR primer pair SEQ ID NO: 145 + 146 from US patent 6,001,564 (Table 24) as well as shorter versions of the original P. aeruginosa-specific primer pair SEQ ID NO: 83 + 84 from US patent 6,001,564 (Table 25). As shown in Tables 24 and 25, both primers of each pair were shortened or lengthen to the same length. Again, those skilled in the art know that the melting temperature of both primers from a pair should be similar to avoid preferential binding at one primer binding site which is

detrimental in PCR (Persing et al., 1993, Diagnostic Molecular Microbiology: Principles and Applications, American Society for Microbiology, Washington, D.C.; Ehrlich and Greenberg, 1994, PCR-based Diagnostics in Infectious Disease, Blackwell Scientific Publications, Boston, MA). All of these shorter or longer primer versions were tested in PCR assays and the efficiency of the amplification was compared with the original primer pair SEQ ID NOs 145 and 146.

Materials and methods

See the Materials and methods section of Example 44.

Results

a) Testing primers with nucleotide changes

The results of the PCR assays with the 13 modified versions of SEQ ID NO: 146 from US patent 6,001,564 are shown in Table 23. The 8 modified primers having a single nucleotide variation showed an efficiency of amplification identical to the original primer pair based on testing with 3 different dilutions of genomic DNA. The four primers having two nucleotide variations and primer VBmut12 having 3 nucleotide changes also showed PCR results identical to those obtained with the original pair. Finally, primer VBmut13 with four nucleotide changes showed a reduction in sensitivity by approximately one log as compared with the original primer pair. However, reducing the annealing temperature from 55 to 50 °C gave an efficiency of amplification very similar to that observed with the original primer pair (Table 23). In fact, reducing the annealing temperature of PCR cycles represents an effective way to reduce the stringency of hybridization for the primers and consequently allows the binding of probes with mismatches (Persing et al., 1993, Diagnostic Molecular Microbiology: Principles and Applications, American Society for Microbiology, Washington, D.C.). Subsequently, we have confirmed the

specificity of the PCR assays with each of these 13 modified versions of SEQ ID NO: 146 from US patent 6,001,564 by performing amplifications from all bacterial species closely related to *S. epidermidis* which are listed in Table 22.

b) Testing shorter or longer versions of primers

For these experiments, two primer pairs were selected: i) SEQ ID NO: 145 + 146 from US patent 6,001,564 (specific to *S. epidermidis*) which are AT rich and ii) SEQ ID NO: 83 + 84 (specific to *P. aeruginosa*) which are GC rich. For the AT rich sequence, primers of 15 to 30 nucleotide in length were designed (Table 24) while for the GC rich sequences, primers of 13 to 19 nucleotide in length were designed (Table 25).

Table 24 shows that, for an annealing temperature of 55 °C, the 30-25-, 20- and 17-nucleotide versions of SEQ ID NO: 145 and 146 from US patent 6,001,564 all showed identical results as compared with the original primer pair except that the 17-nucleotide version amplified slightly less efficiently the *S. epidermidis* DNA. Reducing the annealing temperature from 55 to 45 °C for the 17-nucleotide version allowed to increase the amplification efficiency to a level very similar to that with the original primer pair (SEQ ID NO: 145 + 146 from US patent 6,001,564). Regarding the 15-nucleotide version, there was amplification of *S. epidermidis* DNA only when the annealing temperature was reduced to 45 °C. Under those PCR conditions the assay remained *S. epidermidis*-specific but the amplification signal with *S. epidermidis* DNA was sligthly lower as compared with the original primer pair. Subsequently, we have further confirmed the specificity of the shorter or longer versions by amplifying DNA from all bacterial species closely related to *S. epidermidis* which are listed in Table 22.

Table 25 shows that, for an annealing temperature of 55 °C, all shorter versions of SEQ ID NO: 83 and 84 from US patent 6,001,564 showed identical PCR results as

compared with the original primer pair. As expected, these results show that it is simpler to reduce the length of GC rich as compared with AT rich. This is attributable to the fact that GC binding is more stable than AT binding.

Conclusion

a) Testing primers with nucleotide changes

The above experiments clearly show that PCR primers may be modified at one or more nucleotide positions without affecting the specificity and the sensitivity of the PCR assay. These results strongly suggest that a given oligonucleotide can detect variant genomic sequences from the target species. In fact, the nucleotide changes in the selected primers were purposely introduced at the third position of each codon to mimic nucleotide variation in genomic DNA. Thus we conclude that it is justified to claim "a variant thereof" for i) the SEQ IDs of the fragments and oligonucleotides which are object of the present patent application and ii) genomic variants of the target species.

b) Testing shorter or longer versions of primers

The above experiments clearly show that PCR primers may be shorter or longer without affecting the specificity and the sensitivity of the PCR assay. We have showed that oligonucleotides ranging in sizes from 13 to 30 nucleotides may be as specific and sensitive as the original primer pair from which they were derived. Consequently, these results suggest that it is not exaggerated to claim sequences having at least 12 nucleotide in length.

This invention has been described herein above, and it is readily apparent that modifications can be made thereto without departing from the spirit of this invention. These modifications are under the scope of this invention, as defined in the appended claims.

Table 1. Distribution (%) of nosoc mial path gens for vari us human infections in USA (1990-1992).

5	Pathogen	UTI ²	SSI ³	BSI⁴	Pneumonia	CSF⁵
,	Escherichia coli	27	9	5	4	2
	Staphylococcus aureus	2	21	17	21	2
	Staphylococcus epidermidis	2	6	20	0	1
	Enterococcus faecalis	16	12	9	2	0
10	Enterococcus faecium	1	1	0	0	0
	Pseudomonas aeruginosa	12	9	3	18	0
	Klebsiella pneumoniae	7	3	4	9	0
	Proteus mirabilis	5	3	1	2	0
	Streptococcus pneumoniae	0	0	3	1	18
15	Group B Streptococci	1	1	2	1	6
	Other streptococci	3	5	2	1	3
	Haemophilus influenzae	0	0	0	6	45
	Neisseria meningitidis	0	0	0	0	14
	Listeria monocytogenes	0	0	0	0	3
20	Other enterococci	1	1	0	0	0
	Other staphylococci	2	8	13	2	0
	Candida albicans	9	3	5	5	0
	Other Candida	2	1	3	1	0
	Enterobacter sp.	5	7	4	12	2
25	Acinetobacter sp.	1	1	2	4	2
	Citrobacter sp.	2	1	1	1	0
	Serratia marcescens	1	1	1	3	1
	Other Klebsiella	1	1	1	2	1
	Others	0	6	4	5	00

³⁰

Data recorded by the National Nosocomial Infections Surveillance (NNIS) from 80 hospitals (Emori and Gaynes, 1993, Clin. Microbiol. Rev., 6:428-442).

Urinary tract infection.

Surgical site infection.

^{35 &}lt;sup>4</sup> Bloodstream infection.

Cerebrospinal fluid.

Tabl 2. Distribution (%) f bloodstr am infecti n pathogens in Qu bec (1995), Canada (1992), UK (1969-1988) and USA (1990-1992).

	Organism	Quebec	Canada ²	UK ³		USA⁴
				Community- acquired	Hospital- acquired	Hospital- acquired
	E. coli	15.6	53.8	24.8	20.3	5.0
)	S. epidermidis and other CoNS	25.8	-	0.5	7.2	31.0
	S. aureus	9.6	-	9.7	19.4	16.0
	S. pneumoniae	6.3	-	22.5	2.2	-
	E. faecalis	3.0	-	1.0	4.2	-
5	E. faecium	2.6	-	0.2	0.5	-
	Enterococcus sp.	-	-		9.0	
	H. influenzae	1.5	-	3.4	0.4	-
	P. aeruginosa	1.5	8.2	1.0	8.2	3.0
	K. pneumoniae	3.0	11.2	3.0	9.2	4.0
0	P. mirabilis	-	3.9	2.8	5.3	1.0
	S. pyogenes	-	-	1.9	0.9	-
	Enterobacter sp.	4.1	5.5	0.5	2.3	4.0
	Candida sp.	8.5	-	-	1.0	8.0
	Others	18.5	17.4	28.7	18.9	19.0

Data obtained for 270 isolates collected at the Centre Hospitalier de l'Université Laval (CHUL) during a 5 month period (May to October 1995).

Data from 10 hospitals throughout Canada representing 941 gram-negative isolates. (Chamberland et al., 1992, Clin. Infect. Dis., 15:615-628).

Data from a 20-year study (1969-1988) for nearly 4000 isolates. (Eykyn et al., 1990, J. Antimicrob. Chemother., Suppl. C, 25:41-58).

Data recorded by the National Nosocomial Infections Surveillance (NNIS) from 80 hospitals (Emori and Gaynes, 1993, *Clin. Microbiol. Rev.*, **6**:428-442).

⁵ Coagulase-negative staphylococci.

Tabl 3. Distributi n f positive and negativ clinical specimens tested at th micr biology laboratory of the CHUL (February 1994 – January 1995).

5	Clinical specimens and/or sites	No. of samples tested (%)	% of positive specimens	% of negative specimens
	Urine	17,981 (54.5)	19.4	80.6
	Blood culture/marrow	10,010 (30.4)	6.9	93.1
	Sputum	1,266 (3.8)	68.4	31.6
10	Superficial pus	1,136 (3.5)	72.3	27.7
	Cerebrospinal fluid	553 (1.7)	1.0	99.0
	Synovial fluid	523 (1.6)	2.7	97.3
	Respiratory tract	502 (1.5)	56.6	43.4
	Deep pus	473 (1.4)	56.8	43.2
15	Ears	289 (0.9)	47.1	52.9
	Pleural and pericardial fluid	132 (0.4)	1.0	99.0
	Peritoneal fluid	101(0.3)	28.6	71.4
	Total:	32,966 (100.0)	20.0	80.0

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Table 4.

Example of microbial species for which tuf and/or atpD and/or recA nucleic acids and/or sequences are used in the present invention.

5	Bacte	Bacterial species					
	Abiotrophia adiacens		Brevibacterium flavum				
	Abiotrophia defectiva		Brevundimonas diminuta				
	Achromobacter xylosoxidans subsp. denitrificans	65	Buchnera aphidicola				
10	Acetobacterium woodi		Budvicia aquatica				
	Acetobacter aceti		Burkholderia cepacia				
	Acetobacter altoacetigenes		Burkholderia mallei				
	Acetobacter polyoxogenes		Burkholderia pseudomallei 🖢				
	Acholeplasma laidlawii	70	Buttiauxella agrestis				
15	Acidothermus cellulolyticus		Butyrivibrio fibrisolvens				
	Acidiphilum facilis		Campylobacter coli				
	Acinetobacter baumannii		Campylobacter curvus				
	Acinetobacter calcoaceticus		Campylobacter fetus subsp. fetus				
	Acinetobacter lwoffii	75	Campylobacter fetus subsp. venerealis				
20	Actinomyces meyeri		Campylobacter gracilis				
	Aerococcus viridans		Campylobacter jejuni				
	Aeromonas hydrophila		Campylobacter jejuni subsp. doylei				
	Aeromonas salmonicida		Campylobacter jejuni subsp. jejuni				
	Agrobacterium radiobacter	80	Campylobacter lari				
25	Agrobacterium tumefaciens		Campylobacter rectus				
	Alcaligenes faecalis subsp. faecalis		Campylobacter sputorum subsp. sputorum				
	Allochromatium vinosum		Campylobacter upsaliensis				
	Anabaena variabilis	0.5	Cedecea davisae				
	Anacystis nidulans	85	Cedecea lapagei				
30	Anaerorhabdus furcosus		Cedecea neteri				
	Aquifex aeolicus		Chlamydia pneumoniae				
	Aquifex pyrophilus		Chlamydia psittaci				
	Arcanobacterium haemolyticum	00	Chlamydia trachomatis				
	Archaeoglobus fulgidus	90	Chlorobium vibrioforme				
35	Azotobacter vinelandii		Chloroflexus aurantiacus				
	Bacillus anthracis		Chryseobacterium meningosepticum				
	Bacillus cereus		Citrobacter amalonaticus				
	Bacillus firmus	95	Citrobacter braakii				
40	Bacillus halodurans	93	Citrobacter farmeri Citrobacter freundii				
40	Bacillus megaterium		Citrobacter koseri				
	Bacillus mycoides		Citrobacter koseri Citrobacter sedlakii				
	Bacillus pseudomycoides		Citrobacter werkmanii				
	Bacillus stearothermophilus Bacillus subtilis	100	Citrobacter youngae				
45	_	100	Clostridium acetobutylicum				
43	Bacillus thuringiensis Pacillus weihenstenhanensis		Clostridium beijerinckii				
	Bacillus weihenstephanensis Bacteroides distasonis		Clostridium bifermentans				
	Bacteroides fragilis		Clostridium botulinum				
	Bacteroides forsythus	105	Clostridium difficile				
50	Bacteroides ovatus		Clostridium innocuum				
50	Bacteroides vulgatus		Clostridium histolyticum				
	Bartonella henselae		Clostridium novyi				
	Bifidobacterium adolescentis		Clostridium septicum				
	Bifidobacterium breve	110	Clostridium perfringens				
55	Bifidobacterium dentium		Clostridium ramosum				
-	Bifidobacterium longum	•	Clostridium sordellii				
	Blastochloris viridis		Clostridium tertium				
	Borrelia burgdorferi		Clostridium tetani				
	Bordetella pertussis	115	Comamonas acidovorans				
60	Bordetella bronchiseptica		Corynebacterium accolens				
	Brucella abortus		Corynebacterium bovis				
	Brevibacterium linens		Corynebacterium cervicis				

Table 4. Example of microbial species for which *tuf* and/or *atpD* and/or *recA* nucleic acids and/or sequences are used in the present invention (c ntinued).

-	Bacter	rial species (c	ontinued)
5	Complete Andrew Higher		Eukastarium lantum
	Corynebacterium diphtheriae	65	Eubacterium lentum Eubacterium nodatum
	Corynebacterium flavescens	03	Ewingella americana
	Corynebacterium genitalium Corynebacterium glutamicum		Francisella tularensis
10	Corynebacterium giuiamicum Corynebacterium jeikeium		Frankia alni
10	Corynebacterium kutscheri		Fervidobacterium islandicum
	Corynebacterium minutissimum	70	Fibrobacter succinogenes
	Corynebacterium mycetoides	, ,	Flavobacterium ferrigeneum
	Corynebacterium pseudodiphtheriticum		Flexistipes sinusarabici
15	Corynebacterium pseudogenitalium		Fusobacterium gonidiaformans
15	Corynebacterium pseudotuberculosis		Fusobacterium necrophorum subsp. necrophorum
	Corynebacterium renale	75	Fusobacterium nucleatum subsp. polymorphum
	Corynebacterium striatum		Gardnerella vaginalis
	Corynebacterium ulcerans		Gemella haemolysans
20	Corynebacterium urealyticum		Gemella morbillorum
_,	Corynebacterium xerosis		Globicatella sanguis
	Coxiella burnetii	80	Gloeobacter violaceus
	Cytophaga lytica		Gloeothece sp.
	Deinococcus radiodurans		Gluconobacter oxydans
25	Deinonema sp.		Haemophilus actinomycetemcomitans
	Edwardsiella hoshinae		Haemophilus aphrophilus
	Edwardsiella tarda	85	Haemophilus ducreyi
	Ehrlichia canis		Haemophilus haemolyticus
	Ehrlichia risticii		Haemophilus influenzae
30	Eikenella corrodens		Haemophilus parahaemolyticus
	Enterobacter aerogenes		Haemophilus parainfluenzae
	Enterobacter agglomerans	90	Haemophilus paraphrophilus
	Enterobacter amnigenus		Haemophilus segnis
	Enterobacter asburiae		Hafnia alvei
35	Enterobacter cancerogenus		Halobacterium marismortui
	Enterobacter cloacae	05	Halobacterium salinarum
	Enterobacter gergoviae	95	Haloferax volcanii
	Enterobacter hormaechei		Helicobacter pylori
40	Enterobacter sakazakii		Herpetoshiphon aurantiacus
40	Enterococcus avium		Kingella kingae
	Enterococcus casseliflavus	100	Klebsiella ornithinolytica
	Enterococcus cecorum	100	Klebsiella oxytoca Klebsiella planticola
	Enterococcus columbae		Klebsiella pneumoniae subsp. ozaenae
45	Enterococcus dispar Enterococcus durans		Klebsiella pneumoniae subsp. pneumoniae
43			
	Enterococcus faecalis Enterococcus faecium	105	Klebsiella pneumoniae subsp. rhinoscleromatis
	Enterococcus flavescens	103	Klebsiella terrigena
	Enterococcus gallinarum		Kluyvera ascorbata
50	Enterococcus hirae		Kluyvera cryocrescens
50	Enterococcus malodoratus		Kluyvera georgiana
	Enterococcus mundtii	110	Kocuria kristinae
	Enterococcus pseudoavium		Lactobacillus acidophilus
	Enterococcus raffinosus		Lactobacillus garvieae
55	Enterococcus saccharolyticus		Lactobacillus paracasei
	Enterococcus solitarius		Lactobacillus casei subsp. casei
	Enterococcus sulfureus	115	Lactococcus garvieae
	Erwinia amylovora		Lactococcus lactis
	Erwinia carotovora		Lactococcus lactis subsp. lactis
60	Escherichia coli		Leclercia adecarboxylata
	Escherichia fergusonii		Legionella micdadei
	Escherichia hermannii		
	Escherichia vulneris		

Table 4.

Example f microbial species for which tuf and/or atpD and/or recA nucleic acids and/or sequences are used in the present inventin (c ntinued).

5	Bacterial species (continued)						
	Legionella pneumophila subsp. pneumophila		Neisseria gonorrhoeae				
	Leminorella grimontii		Neisseria lactamica				
	Leminorella richardii	65	Neisseria meningitidis				
10	Leptospira biflexa		Neisseria mucosa				
	Leptospira interrogans		Neisseria perflava				
	Leuconostoc mesenteroides subsp.		Neisseria pharyngis var. flava				
	dextranicum		Neisseria polysaccharea				
	Listeria innocua	70	Neisseria sicca				
15	Listeria ivanovii		Neisseria subflava				
	Listeria monocytogenes		Neisseria weaveri				
	Listeria seeligeri		Obesumbacterium proteus				
	Macrococcus caseolyticus	0_	Ochrobactrum anthropi				
	Magnetospirillum magnetotacticum	75	Pantoea agglomerans				
20	Megamonas hypermegale		Pantoea dispersa				
	Methanobacterium thermoautotrophicum		Paracoccus denitrificans				
	Methanococcus jannaschii		Pasteurella multocida				
	Methanococcus vannielii	00	Pectinatus frisingensis				
	Methanosarcina barkeri	80	Peptococcus niger				
25	Methanosarcina jannaschii		Peptostreptococcus anaerobius				
	Methylobacillus flagellatum		Peptostreptococcus asaccharolyticus				
	Methylomonas clara		Peptostreptococcus prevotii				
	Micrococcus luteus	05	Phormidium ectocarpi				
20	Micrococcus lylae	85	Pirellula marina				
30	Mitsuokella multacidus		Planobispora rosea				
	Mobiluncus curtisii subsp. holmesii		Plesiomonas shigelloides				
	Moellerella thermoacetica		Plectonema boryanum Porphyromonas asaccharolytica				
	Moellerella wisconsensis	90	Porphyromonas gingivalis				
35	Moorella thermoacetica	90	Pragia fontium				
33	Moraxella catarrhalis		Prevotella buccalis				
	Moraxella osloensis Moracella moragnii subsp. moragnii		Prevotetta baccans Prevotella melaninogenica				
	Morganella morganii subsp. morganii		Prevotella oralis				
	Mycobacterium avium Mycobacterium bovis	95	Prevotella ruminocola				
40	Mycobacterium gordonae	, ,	Prochlorothrix hollandica				
70	Mycobacterium kansasii		Propionibacterium acnes				
	Mycobacterium leprae		Propionigenium modestum				
	Mycobacterium terrae		Proteus mirabilis				
	Mycobacterium tuberculosis	100	Proteus penneri				
45	Mycoplasma capricolum		Proteus vulgaris				
	Mycoplasma gallisepticum		Providencia alcalifaciens				
	Mycoplasma genitalium		Providencia rettgeri				
	Mycoplasma hominis		Providencia rustigianii				
	Mycoplasma pirum	105	Providencia stuartii				
50	Mycoplasma mycoides		Pseudomonas aeruginosa				
	Mycoplasma pneumoniae		Pseudomonas fluorescens				
	Mycoplasma pulmonis		Pseudomonas putida				
	Mycoplasma salivarium	110	Pseudomonas stutzeri				
	Myxococcus xanthus	110	Psychrobacter phenylpyruvicum				
55	Neisseria animalis		Pyrococcus abyssi				
	Neisseria canis		Rahnella aquatilis Rickettsia prowazekii				
	Neisseria cinerea						
	Neisseria cuniculi	115	Rhizobium leguminosarum Rhizobium phaseoli				
40	Neisseria elongata subsp. elongata	113	Rhodobacter capsulatus				
60	Neisseria elongata subsp. intermedia		Rhodobacter capsulatus Rhodobacter sphaeroides				
	Neisseria flava Neisseria flavescens						
	iveisseim juivesceib		_				

Table 4. Example of microbial species f r which tuf and/or atpD and/or recA nucleic acids and/or sequences are used in the present invention (continued).

5

Bacterial species (continued)

	Rhodopseudomonas palustris		Streptococcus gordonii
	Rhodospirillum rubrum	65	Streptococcus macacae
10	Ruminococcus albus		Streptococcus mitis
10	Ruminococcus bromii		Streptococcus mutans
	Salmonella bongori		Streptococcus oralis
	Salmonella choleraesuis subsp. arizonae		Streptococcus parasanguinis
	Salmonella choleraesuis subsp	70	Streptococcus pneumoniae
15	choleraesuis		Streptococcus pyogenes
13	Salmonella choleraesuis subsp.		Streptococcus ratti
	diarizonae		Streptococcus salivarius
	Salmonella choleraesuis subsp.		Streptococcus salivarius subsp. thermophilus
	houtenae	75	Streptococcus sanguinis
20	Salmonella choleraesuis subsp. indica	. •	Streptococcus sobrinus
20	Salmonella choleraesuis subsp. salamae		Streptococcus suis
	Serpulina hyodysenteriae		Streptococcus uberis
			Streptococcus vestibularis
	Serratia ficaria	80	Streptomyces anbofaciens
25	Serratia fonticola	00	Streptomyces aureofaciens
23	Serratia grimesii		Streptomyces cinnamoneus
	Serratia liquefaciens Serratia marcescens		Streptomyces coelicolor
			Streptomyces collinus
	Serratia odorifera	85	Streptomyces lividans
20	Serratia plymuthica	0.5	Streptomyces netropsis
30	Serratia rubidaea		Streptomyces ramocissimus
	Shewanella putrefaciens		Streptomyces rimosus
	Shigella boydii		Streptomyces venezuelae
	Shigella dysenteriae	90	Succinivibrio dextrinosolvens
25	Shigella flexneri	70	Synechococcus sp.
35	Shigella sonnei		Synechocystis sp.
	Sinorhizobium meliloti		Tatumella ptyseos
	Spirochaeta aurantia		Taxeobacter occealus
	Staphylococcus aureus	95	Tetragenococcus halophilus
40	Staphylococcus aureus subsp. aureus),	Thermoplasma acidophilum
40	Staphylococcus auricularis		Thermotoga maritima
	Staphylococcus capitis subsp. capitis		Thermus aquaticus
	Staphylococcus cohnii subsp. cohnii		Thermus thermophilus
	Staphylococcus epidermidis	100	Thiobacillus ferrooxidans
45	Staphylococcus haemolyticus	100	Thiomonas cuprina
45	Staphylococcus hominis		Trabulsiella guamensis
	Staphylococcus hominis subsp. hominis		Treponema pallidum
	Staphylococcus lugdunensis		Ureaplasma urealyticum
	Staphylococcus saprophyticus	105	Veillonella parvula
50	Staphylococcus sciuri subsp. sciuri	103	Vibrio alginolyticus
50	Staphylococcus simulans		Vibrio anguillarum
	Staphylococcus warneri		Vibrio cholerae
	Stigmatella aurantiaca		Vibrio mimicus
	Stenotrophomonas maltophilia	110	Wolinella succinogenes
	Streptococcus acidominimus	110	Xanthomonas citri
55	Streptococcus agalactiae		Xanthomonas oryzae
	Streptococcus anginosus		Xenorhabdus bovieni
	Streptococcus bovis		Xenorhabdus nematophilus
	Streptococcus cricetus	115	Yersinia bercovieri
<i>(</i> 0	Streptococcus cristatus	113	Yersinia enterocolitica
60	Streptococcus downei		Yersinia frederiksensii
	Streptococcus dysgalactiae		Yersinia intermedia
	Streptococcus equi subsp. equi		Yersinia pestis
	Streptococcus ferus		

Table 4. Example of microbial species f r which tuf and/or atpD and/or recA nucleic acids and/or sequences are used in the present invention (continued).

Bacterial species (continued)

Yersinia pseudotuberculosis Yersinia rohdei Yokenella regensburgei 200gloea ramigera

5

Table 4. Example of micr bial species f r which tuf and/or atpD and/ r recA nucleic acids and/or sequences are used in the present inventin (c ntinued).

5		Fungal speci	ies
	Absidia corymbifera		Fusarium moniliforme
	Absidia glauca	(5	Fusarium oxysporum
• •	Alternaria alternata	65	Fusarium solani
10	Arxula adeninivorans		Geotrichum sp.
	Aspergillus flavus		Histoplasma capsulatum
	Aspergillus fumigatus		Hortaea werneckii
	Aspergillus nidulans	70	Issatchenkia orientalis Kudrjanzev
	Aspergillus niger	70	Kluyveromyces lactis
15	Aspergillus oryzae		Malassezia furfur
	Aspergillus terreus		Malassezia pachydermatis
	Aspergillus versicolor		Malbranchea filamentosa
	Aureobasidium pullulans	25	Metschnikowia pulcherrima
	Basidiobolus ranarum	75	Microsporum audouinii
20	Bipolaris hawaiiensis		Microsporum canis
	Bilophila wadsworthia		Mucor circinelloides
	Blastoschizomyces capitatus		Neurospora crassa
	Blastomyces dermatitidis	00	Paecilomyces lilacinus
	Candida albicans	80	Paracoccidioides brasiliensis
25	Candida catenulata		Penicillium marneffei
	Candida dubliniensis		Phialaphora verrucosa
	Candida famata		Pichia anomala
	Candida glabrata	0.5	Piedraia hortai
	Candida guilliermondii	85	Podospora anserina
30	Candida haemulonii		Podospora curvicolla
	Candida inconspicua		Puccinia graminis
	Candida kefyr		Pseudallescheria boydii
	Candida krusei	00	Reclinomonas americana
	Candida lambica	90	Rhizomucor racemosus
35	Candida lusitaniae		Rhizopus oryzae
	Candida norvegica		Rhodotorula minuta
	Candida norvegensis		Rhodotorula mucilaginosa
	Candida parapsilosis	0.5	Saccharomyces cerevisiae
4.0	Candida rugosa	95	Saksenaea vasiformis
40	Candida sphaerica		Schizosaccharomyces pombe
	Candida tropicalis		Scopulariopsis koningii
	Candida utilis		Sordaria macrospora
	Candida viswanathii	100	Sporobolomyces salmonicolor
4.5	Candida zeylanoides	100	Sporothrix schenckii
45	Cladophialophora carrionii		Stephanoascus ciferrii
	Coccidioides immitis		Syncephalastrum racemosum
	Coprinus cinereus		Trichoderma reesei Trichophyton mentagrophytes
	Cryptococcus albidus	105	Trichophyton rubrum
50	Cryptococcus humicolus	103	Trichophyton tonsurans
50	Cryptococcus laurentii		Trichosporon cutaneum
	Cryptococcus neoformans		Ustilago maydis
	Cunninghamella bertholletiae		Wangiella dermatitidis
	Curvularia lunata	110	Yarrowia lipolytica
55	Emericella nidulans	110	ταπονία προιγίτα
33	Emmonsia parva		
	Eremothecium gossypii		
	Exophiala dermatitidis		
	Exophiala jeanselmei		
60	Exophiala moniliae		
ou	Exserohilum rostratum Examothacium gossynii		
	Eremothecium gossypii		
	Fonsecaea pedrosoi		

Table 4. Example of micr bial species f r which tuf and/ r atpD and/or recA nucleic acids and/or sequences are used in the present invention (continued).

Parasitical species
Dah saia hisamina
Babesia bigemina
Babesia bovis
Babesia microti
Blastocystis hominis
Crithidia fasciculata
Cryptosporidium parvum
Entamoeba histolytica
Giardia lamblia
Kentrophoros sp.
Leishmania aethiopica
Leishmania amazonensis
Leishmania braziliensis
Leishmania donovani
Leishmania infantum
Leishmania enriettii
Leishmania gerbilli
Leishmania guyanensis
Leishmania hertigi
Leishmania major
Leishmania mexicana
Leishmania panamensis
Leishmania tarentolae
Leishmania tropica
Neospora caninum
Onchocerca volvulus
Plasmodium berghei
Plasmodium falciparum
Plasmodium knowlesi
Porphyra purpurea
Toxoplasma gondii
Treponema pallidum
Trichomonas tenax
Trichomonas vaginalis
Trypanosoma brucei
Trypanosoma brucei subsp. brucei
Trypanosoma congolense
Trypanosoma cruzi

Table 5. Antimicrobial agents resistance genes selected for diagn stic purposes.

Gene	Antimicrobial agent	Bacteria ¹	ACCESSION NO.	SEQ ID NO
aac(3)-lb ²	Aminoglycosides	Enterobacteriaceae Pseudomonads	L06157	
aac(3)-IIb ²	Aminoglycosides	Enterobacteriaceae, Pseudomonads	M97172	
aac(3)-IVa 2	Aminoglycosides	Enterobacteriaceae	X01385	
aac(3)-IVa ² aac(3)-VIa ²	Aminoglycosides	Enterobacteriaceae,	M88012	
		Pseudomonads		
aac(2')-1a ²	Aminoglycosides	Enterobacteriaceae, Pseudomonads	X04555	
aac(6')-aph(2'') ²	Aminoglycosides	Enterococcus sp., Staphylococcus sp.		83-86 ³
aac(6')-Ia, ²	Aminoglycosides	Enterobacteriaceae, Pseudomonads	M18967	
aac(6')-Ic ²	Aminoglycosides	Enterobacteriaceae, Pseudomonads	M94066	
aac(6')-IIa ²	Aminoglycosides	Pseudomonads		112 ⁴
aadB [ant(2")-Ia 2]	Aminoglycosides	Enterobacteriaceae		53-54 ³
aacCl [aac(3)-la 2]	Aminoglycosides	Pseudomonads		55-56 ³
$aacC2 [aac(3)-IIa^{2}]$	Aminoglycosides	Pseudomonads		57-58 ³
$aacC3$ [$aac(3)$ -III 2]	Aminoglycosides	Pseudomonads		59-60 ³
aacA4 [aac(6')-Ib 2]	Aminoglycosides	Pseudomonads		65-66 ³
ant(3")-la ²	Aminoglycosides	Enterobacteriaceae.	X02340	35 50
, ,		Enterococcus sp., Staphylococcus sp.	M10241	
ant(4')-Ia ²	Aminoglycosides	Staphylococcus sp.	V01282	
ant(4')-Ia ² aph(3')-Ia ²	Aminoglycosides	Enterobacteriaceae, Pseudomonads	J01839	
aph(3')-IIa ²	Aminoglycosides	Enterobacteriaceae, Pseudomonads	V00618	
aph(3')-IIIa ²	Aminoglycosides	Enterococcus sp., Staphylococcus sp.	V01547	
aph(3')-VIa ²	Aminoglycosides	Enterobacteriaceae, Pseudomonads	X07753	
rpsL ²	Streptomycin	M. tuberculosis,	X80120	
	-	M. avium complex	U14749	
		-	X70995	
			L08011	
blaOXA 5,6	ß-lactams	Enterobacteriaceae,	Y10693	110 4
		Pseudomonads	AJ238349	
			AJ009819	
			X06046	
			X03037	
			X07260	
			U13880	
			X75562	
			AF034958	
			J03427	
			Z22590	
			U59183	
			L38523	
			U63835	
			AF043100	
			AF060206 U85514	
			AF043381	
			AF024602	
			AF064820	45-48 ³
	A 1 .	77 7.71		45 AQ 3
bla _{ROB} 5	B-lactams	Haemophilus sp. Pasteurella sp.		43-40

Table 5. Antimicr bial agents resistance genes selected for diagnostic purposes (c ntinued).

blaSHV ⁵ ,	б ß-lactams	Enterobacteriacea, Pseudomonas aeruginosa	AF124984 AF148850	41-44 3
			M59181 X98099	
			M33655	
			AF148851	
			X53433 L47119	
			AF074954	
			X53817 AF096930	
			X55640	
			Y11069	
			U20270 U92041	
			S82452	
			X98101	
			X98105 AF164577	
			AJ011428	
			AF116855	
	•		AB023477 AF293345	
			AF227204	
			AF208796	
blames 45	,6 ß-lactams	Enterobacteriaceae,	AF132290 AF012911	37-40 ³
ola TEM	blaTEM 5,6 B-lactams	Neisseria sp.,	U48775	37-40
		Haemophilus sp.	AF093512	
			AF052748 X64523	
			Y13612	
			X57972	
			AF157413 U31280	
			U36911	
			U48775	
			V00613 X97254	
			AJ012256	
			X04515	
			AF126482 U09188	
			M88143	
			Y14574	
			AF188200 AJ251946	
			Y17581	
			Y17582 Y17583	
			M88143	
			U37195	
			Y17584 X64523	
			U95363	
			Y10279	
			Y10280 Y10281	
			AF027199	
			AF104441	
			AF104442 AF062386	
			X57972	
			AF047171	
			AF188199 AF157553	
			AF190694	
			AF190695	
			AF190693 AF190692	

Table 5. Antimicr bial agents resistance genes selected for diagn stic purposes (c ntinued).

Gene	Antimicrobial agent	Bacteria ¹	ACCESSION NO.	SEQ ID NO
bla _{CARB} 5	B-lactams	Pseudomonas sp.,	J05162	
OWCAKB	., 144 Mills	Enterobacteriaceae	S46063	
		2/110/00/00/00/00/00	M69058	
			U14749	
			D86225	
			D13210	
			Z18955	
			AF071555	
			AF153200	
-			AF030945	
bla _{CTX-M-1} 5	B-lactams	Enterobacteriaceae	X92506	
bla _{CTX-M-2} 5	B-lactams	Enterobacteriaceae	X92507	
bla _{CMY-2} 7	B-lactams	Enterobacteriaceae	X91840	
CM1-2		3	AJ007826	
			AJ011293	
			AJ011291	
			Y17716	
			Y16783	
			Y16781	
			Y15130	
			U77414 S83226	
			Y15412	
			X78117	
bla _{IMP} 5	B-lactams	Enterobacteriaceae,	AJ223604	
2.722		Pseudomonas aeruginos	a S71932	
		G	D50438	
			D29636	
			X98393	
			AB010417	
_			D78375	
bla _{PER-1} 5	B-lactams	Enterobacteriaceae,	Z21957	
_		Pseudomodanaceae		
bla _{PER-2} ⁷	ß-lactams	Enterobacteriaceae	X93314	4
blaZ ¹²	B-lactams	Enterococcus sp., Staphylococcus sp.		111 4
mecA ¹²	B-lactams	Staphylococcus sp.		97-98 ³

Table 5. Antimicr bial agents resistance genes selected for diagnostic purp ses (c ntinued).

Gene	Antimicrobial agent	Bacteria ¹	ACCESSION NO.	SEQ ID NO.
pbp1a ¹³	B-lactams	Streptococcus pneumoniae	?	1004-1018,
,opiu			M90527	1648,2056-2064
			X67872	2273-2276
			AB006868	
			AB006874	
			X67873	
			AB006878 AB006875	
			AB006877	
			AB006879	
			AF046237	
			AF046235	
			AF026431	
			AF046232	
			AF046233	
			AF046236 X67871	
			Z49095	
		•	AF046234	
			AB006873	
			X67866	
			X67868	
			AB006870	
			AB006869	
			AB006872	
			X67870 AB006871	
			X67867	
			X67869	
			AB006876	
			AF046230	
			AF046238	
pbp2b.13	ß-lactams	Streptococcus pneumonia	Z49094	1019-1033
pop20:	b-factains	Streptococcus pheumoma	X16022	1019-1033
			M25516	
			M25518	
			M25515	
			U20071	
			U20084	
			U20082	
			U20067	
			U20079 Z22185	
			U20072	
pbp2b 13	B-lactams	Streptococcus pneumonia	e U20083	
<i>pop20</i>	p-lactains	sirepiococcus preumonie	U20081	
			M25522	
			U20075	
			U20070	
			U20077	
			U20068	
			Z22184 U20069	
			U20078	
			M25521	
			M25525	
			M25519	
			Z21981	
			M25523	
			M25526	
			U20076	
			U20074	
			M25520 M25517	
			M25524	
			Z22230	
			U20073	
			U20080	
			020080	

Table 5. Antimicr bial agents resistance genes selected f r diagn stic purposes (c ntinued).

Gene	Antimicrobial agent	Bacteria ¹	ACCESSION NO.	SEQ ID 1
pbp2x 13	B-lactams	Streptococcus pneumoniae	X16367	1034-104
			X65135	
			AB011204	
			AB011209	
			AB011199	
			AB011200	
			AB011201 AB011202	
			AB011202 AB011198	
			AB011208	
			AB011205	
			AB015852	
			AB011210	
			AB015849	
			AB015850	
			AB015851	
			AB015847	
			AB015846	
			AB011207	
			AB015848 Z49096	
int	-lactams,	Enterobacteriaceae,	249090	99-102 ³
	trimethoprim			100 106
sul	aminoglycosides,	Pseudomonads		103-106
	antiseptic,			
ermA 14	chloramphenicol Macrolides,	Stanbulacacous an		113 4
ermA - ·	lincosamides,	Staphylococcus sp.		113
	streptogramin B			
ermB 14	Macrolides,	Enterobacteriaceae,		114 4
CIND	Waciondos,	Staphylococcus sp.		
	lincosamides,	Enterococcus sp.		
	streptogramin B	Streptococcus sp.		
ermC 14	Macrolides,	Enterobacteriaceae,		115 ⁴
	lincosamides,	Staphylococcus sp.		
	streptogramin B			
ereA 12	Macrolides	Enterobacteriaceae,	M11277	
		Staphylococcus sp.	E01199	
n 12	34	Property	AF099140	
ereB 12	Macrolides	Enterobacteriaceae	A15097	
msrA 12	Maggalidas	Staphylococcus sp.	X03988	77-80 ³
	Macrolides	Staphylococcus sp.		11-00
mefA, mefE ⁸	Macrolides	Streptococcus sp.	U70055	
mphA 8	Macrolides	Enterobacteriaceae.	U83667 D16251	
трпл С	iviacrondes	Staphylococcus sp.	U34344	
		supreyiococcus sp.	U36578	
linA/linA ^{,9}	Lincosamides	Staphylococcus sp.	J03947	
		suprifications op.	M14039	
			A15070	
4.0			E01245	
linB 10	Lincosamides	Enterococcus faecium	AF110130 AJ238249	
15				20 20 3
vga 15	Streptrogramin	Staphylococcus sp.	M90056 U82085	89-90 ³
vgb 15	Streptrogramin	Staphylococcus sp.	M36022	
0-			M20219	
			AF015628	

Table 5. Antimicrobial agents resistance genes selected for diagnostic purposes (continued).

Gene	Antimicrobial agent	Bacteria ^I	ACCESSION NO	. SEQ ID NO.
vat 15	Streptrogramin	Staphylococcus sp.	L07778	87-88 3
vatB 15	Streptrogramin	Staphylococcus sp.	U19459 L38809	
satA 15	Streptrogramin	Enterococcus faecium	L12033	81-82 3
тирА 12	Mupirocin	Staphylococcus aureus	X75439	01-02
nupri	Maph com	Diaphylococcus aureus	X59478	
			X59477	
gyrA 16	Quinolones	Gram-positive and	X95718	1255, 1607-1608,
		gram-negative bacteria	X06744	1764-1776,
		•	X57174	2013-2014,
			X16817 X71437	2277-2280
			AF065152	
		•	AF060881	
			D32252	
parC/grlA 16	Quinolones	Gram-positive and	AB005036	1777-1785
Ü	-	gram-negative bacteria	AF056287	
			X95717	
			AF129764	
			AB017811	
			AF065152	
parE/grlB 16	Quinolones	Gram-positive bacteria	X95717	
purzigrib	Quinolones	Oram positive bacteria	AF065153	
			AF058920	
norA 16	Quinolones	Staphylococcus sp.	D90119	
		. ,	M80252	
16			M97169	
mexR (nalB) 16 nfxB 16	Quinolones	Pseudomonas aeruginosa Pseudomonas aeruginosa Gram-positive and gram-negative bacteria	U23763	
cat 12	Quinolones Chloramphenicol		X65646 M55620	
cai			X15100	
			A24651	
			M28717	
			A00568	
			A00569	
			X74948	
			Y00723	
			A24362 A00569	
			M93113	
			M62822	
•			M58516	
			V01277	
			X02166	
			M77169	
			X53796	
			J01841 X07848	
			AU/040	
<i>ppflo-</i> like	Chloramphenicol		AF071555	
emhR 1/	Ethambutol	Mycobacterium tuberculosis	U68480	
pncA 17	Pyrazinamide	Mycobacterium tuberculosis	U59967	
троВ ¹⁷	Rifampin	Mycobacterium tuberculosis	AF055891	
	•	•	AF055892	
			S71246	
			L27989	
			A T30	
inhA 17	Isoniazid	Mycobacterium tuberculosis	AF055893 AF106077	

Table 5. Antimicrobial agents resistance genes selected f r diagn stic purp ses (c ntinued).

	Gene	Antimicrobial agent	Bacteria ¹	ACCESSION NO.	SEQ ID NO.
•	vanA 12	Vancomycin	Enterococcus sp.		67-70 ³
			•		1049-1057
	vanB 12	Vancomycin	Enterococcus sp.		116 ⁴
	vanCI 12	Vancomycin	Enterococcus gallinarum		1174
		•	•		1058-1059
	vanC2 12	Vancomycin	Enterococcus casseliflavus		1060-1063
		•	-	U94521	
				U94522	•
				U94523	
				U94524	
				U94525	
				L29638	
	vanC3 12	Vancomycin	Enterococcus flavescens		1064-1066
			-	L29639	
				U72706	
	vanD 18	Vancomycin	Enterococcus faecium	AF130997	
	vanE 12	Vancomycin	Enterococcus faecium	AF136925	
	tetB 19	Tetracycline	Gram-negative bacteria	J01830	
		•		AF162223	
				AP000342	
				S83213	
				U81141	
				V00611	
	tetM 19	Tetracycline	Gram-negative and	X52632	
			Gram-positive bacteria	AF116348	
				U50983	
				X92947	
				M211136	
				U08812	
	20			X04388	
sul	sul II ²⁰	Sulfonamides	Gram-negative bacteria	M36657	
				AF017389	
	20			AF017391	
	dhfrla ²⁰	Trimethoprim	Gram-negative bacteria	AJ238350	
				X17477	
				K00052	
				U09476	
	20	m :		X00926	
	dhfrIb ²⁰	Trimethoprim	Gram-negative bacteria	Z50805	
		motor at an alm	C bostonia	Z50804	
	dhfrV 20	Trimethoprim	Gram-negative bacteria	X12868 Z86002	
	dhfrVI 20	Trimethoprim	Gram-negative bacteria		
	dhfrVII 20	Trimethoprim	Gram-negative bacteria	U31119	
				AF139109	
	dhfrVIII ²⁰	Trimathania	Gram nagativa hastoria	X58425 U10186	
	anjrviii 20	Trimethoprim	Gram-negative bacteria	U09273	
	dhfrIX 20	Trimotharrim	Gram-negative bacteria	X57730	
	dhfrXII 20	Trimethoprim Trimethoprim	Gram-negative bacteria	Z21672	
	aiyi Ali ==	Trimethoprim	Grain-negative vacteria	AF175203	
				AF180731	
				M84522	
	dhfrXIII 20	Trimethoprim	Gram-negative bacteria	Z50802	
	dhfrXV 20	Trimethoprim Trimethoprim	Gram-negative bacteria	Z83331	
	dhfrXVII 20		Gram-negative bacteria	AF170088	
	wyravu	Trimethoprim	Orani-negative vaciena	AF180469	
				AF169041	
				VI.103041	

PCT/CA00/01150 WO 01/23604

Table 5. Antimicrobial agents resistanc genes selected for diagn stic purposes (continued).

Gene	Antimicrobial agent	Bacteria ¹	ACCESSION NO. SEQ ID NO.
5 dfrA 20	Trimethoprim	Staphylococcus sp.	AF045472
3	•		U40259
			AF051916
			X13290
0			Y07536
			Z16422
			Z48233

15 Bacteria having high incidence for the specified antibiotic resistance gene. The presence of the 1 antibiotic resistance genes in other bacteria is not excluded.

Shaw, K. J., P. N. Rather, R. S. Hare, and G. H. Miller. 1993. Molecular genetics of aminoglycoside resistance genes and familial relationships of the aminoglycoside-modifying enzymes. Microbiol. Rev. 57:138-163.

Antibiotic resistance genes from our assigned US patent no. 6,001,564 for which we have selected PCR primer pairs.

These SEQ ID NOs. refer to a previous patent (publication WO98/20157).

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Bush, K., G.A. Jacoby and A. Medeiros. 1995. A functional classification scheme for ß-lactamase and its correlation with molecular structure. Antimicrob. Agents. Chemother. 39:1211-1233. 25

Nucleotide mutations in blaSHV, blaTEM, and blaOXA, are associated with extended-spectrum ßlactamase or inhibitor-resistant ß-lactamase.

Bauerfeind, A., Y. Chong, and K. Lee. 1998. Plasmid-encoded AmpC beta-lactamases: how far have we gone 10 ears after discovery? Yonsei Med. J. 39:520-525.

Sutcliffe, J., T. Grebe, A. Tait-Kamradt, and L. Wondrack. 1996. Detection of erythromycin-resistant 30 determinants by PCR. Antimicrob. Agent Chemother. 40:2562-2566.

Leclerc, R., A., Brisson-Noël, J. Duval, and P. Courvalin. 1991. Phenotypic expression and genetic heterogeneity of lincosamide inactivation in Staphylococcus sp. Antimicrob. Agents. Chemother. 31:1887-1891.

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11 Cockerill III, F.R. 1999. Genetic methods for assessing antimicrobial resistance. Antimicrob. Agents. Chemother. 43:199-212

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13 Dowson, C. G., T. J. Tracey, and B. G. Spratt. 1994. Origin and molecular epidemiology of penicillin-binding-protein-mediated resistance to ß-lactam antibiotics. Trends Molec. Microbiol.2: 361-366.

14 Jensen, L. B., N. Frimodt-Moller, F. M. Aarestrup. 1999. Presence of erm gene classes in Gram-

positive bacteria of animal and human origin in Denmark. FEMS Microbiol. 170:151-158.

Thal, L. A., and M. J. Zervos. 1999. Occurrence and epidemiology of resistance to virginimycin and streptrogramins. J. Antimicrob. Chemother. 43:171-176-

16 Martinez J. L., A. Alonso, J. M. Gomez-Gomez, and F. Baquero. 1998. Quinolone resistance by 50 mutations in chromosomal gyrase genes. Just the tip of the iceberg? J. Antimicrob. Chemother. 42:683-688

17 Cockerill III, F.R. 1999. Genetic methods for assessing antimicrobial resistance. Antimicrob. Agents. Chemother. 43:199-212.

18 Casadewall, B. and P. Courvalin. 1999 Characterization of the vanD glycopeptide resistance gene 55 cluster from Enterococcus faecium BM 4339. J. Bacteriol. 181:3644-3648.

19 Roberts, M.C. 1999. Genetic mobility and distribution of tetracycline resistance determinants. Ciba Found. Symp. 207:206-222.

20 Huovinen, P., L. Sundström, G. Swedberg, and O. Sköld. 1995. Trimethoprim and sulfonamide resistance. Antimicrob. Agent Chemother. 39:279-289.

Table 6. List of bacterial t xins select d for diagn stic purposes.

	Organism	Toxin	Accessi n number
	Actinobacillus actinomycetemcomitans	Cytolethal distending toxin (cdtA, cdtB, cdtC)	AF006830
	Actinomyces pyogenes Aeromonas hydrophila	Leukotoxin (ltxA) Hemolysin (pyolysin) Aerolysin (aerA)	M27399 U84782 M16495
)		Haemolysin (hlyA)	U81555
5	Bacillus anthracis Bacillus cereus	Cytotonic enterotoxin (alt) Anthrax toxin (cya) Enterotoxin (bceT)	L77573 M23179 D17312 AF192766, AF192767
		Enterotoxic hemolysin BL	AJ237785
)	Bacillus mycoides Bacillus pseudomycoides Bacteroides fragilis	Non-haemolytic enterotoxins A,B and C (nhe) Hemolytic enterotoxin HBL Hemolytic enterotoxin HBL Enterotoxin (bftP)	Y19005 AJ243150 to AJ243153 AJ243154 to AJ243156 U67735
5		Matrix metalloprotease/enterotoxin (fragilysin)	S75941, AF038459
		Metalloprotease toxin-2	U90931 AF081785
)	Bordetella bronchiseptica	Metalloprotease toxin-3 Adenylate cyclase hemolysin (cyaA)	AF056297 Z37112, U22953
		Dermonecrotic toxin (dnt)	U59687 AB020025
5	Bordetella pertussis	Pertussis toxin (S1 subunit, tox)	AJ006151 AJ006153 AJ006155 AJ006157 AJ006159
)			AJ007363 M14378, M16494 AJ007364 M13223 X16347
5		Adenyl cyclase (cya)	18323
0	Campylobacter jejuni Citrobacter freundii Clostridium botulinum	Dermonecrotic toxin (dnt) Cytolethal distending toxin (cdtA, cdtB, cdtC) Shiga-like toxin (slt-IIcA) Botulism toxin (BoNT) (A,B,E and F serotypes are neurotoxic for humans; the other serotypes have not been considered)	U10527 U51121 X67514, S53206 X52066, X52088 X73423 M30196
5			X70814 X70819 X71343 Z11934
0			X70817 M81186 X70818 X70815 X62089
5			X62683 S76749 X81714 X70816

Table 6. List of bacterial toxins selected f r diagn stic purposes (continued).

_	Organism	Toxin	Accession number
,	Clostridium botulinum (continued)	•	X70820
•	Clostriatum Votatinum (continuca)		X70281
			L35496
			M92906
-	Clostridium difficile	A toxin (enterotoxin) (tcdA) (cdtA)	AB012304
•	Ciosiriatum atfficite	Te tokin (chierotokin) (teart) (tan)	AF053400
			Y12616
			X51797
			X17194
			M30307
		B toxin (cytotoxin) (toxB) (cdtB)	Z23277
		b tour (e) totolini, (toub)	X53138
,	Clostridium perfringens	Alpha (phospholipase C) (cpa)	L43545
•	Ciosiriaiani perjringera	Auphia (phosphonpase C) (opa)	L43546
			L43547
			L43548
			X13608
			X17300
			D10248
		Beta (dermonecrotic protein) (cpb)	L13198
			X83275
			L77965
	•	5	A 1000766
		Enterotoxin (cpe)	AJ000766
			M98037
			X81849
			X71844
			Y16009
		Enterotoxin pseudogene (not expressed)	AF037328
		Enterotoxin pseudogene (not expresses)	AF037329
			AF037330
		Epsilon toxin $(etxD)$	M80837
			M95206
			X60694
		Iota (Ia and Ib)	X73562
		Lambda (metalloprotease)	D45904
		Theta (parfringalysis (1)	M36704
	Classidiana and alli	Theta (perfringolysin O) Cytotoxin L	X82638
	Clostridium sordellii		X06214
	Clostridium tetani	Tetanos toxin	X04436
	Corynebacterium diphtheriae	Diphtheriae toxin	X00703
		Phospholipase C	A21336
	Corynebacterium pseudotuberculosis	r nosphonpase C	ALIJJU
	Eikenella corrodens	lysine decarboxylase (cadA)	U89166
	Enterobacter cloacae	Shiga-like toxin II	Z50754, U33502
	Enterococcus faecalis	Cytolysin B (cylB)	M38052
	Escherichia coli (EHEC)	Hemolysin toxin (hlyA and ehxA)	AF043471
	Dominiona con (Direct)	, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	X94129
			X79839
			X86087
			AB011549
			AF074613

Table 6. List f bacterial toxins selected for diagnostic purp ses (continued).

_ (Organism	T xin	Accessi n number
_	Escherichia coli (EHEC)	Shiga-like (Vero cytotoxin) (stx)	X81418, M36727
4	Escherichia con (EREC)	Singa-like (Velo Cytotoxiii) (Six)	M14107, E03962
		;	M10133, E03959
			M12863, X07865
			X81417, Y10775
			X81416, Z50754
			X81415, X67515
			Z36900, AF04362
			L11078, M19473
			L04539, M17358
			L11079, M19437
			X65949, M24352
			M21534, X07903
			M29153, Z36899
			Z37725
			Z36901
			X61283
			AB017524
			U72191
			X61283
	Escherichia coli (ETEC)	Enterotoxin (heat-labile) (eltB)	M17874
	Louise to the (Like)		M17873
			J01605
			AB011677
		Enterotoxin (heat-stable) (astA) (estA1)	L11241
		Ditorotomi (non billion) (non) (con e)	M58746
			M29255
			V00612
			J01831
		Control of distanting town	U03293
	Escherichia coli (other)	Cytolethal-distending toxin	
		(cdt) (3 genes)	U04208 U89305
		Cytotoxic necrotizing factor 1 (cnf1)	U42629
		Microsin 24 (mtF)	U47048
		Microcin 24 (mtfS)	AF056581
		Autotransporter enterotoxin (Pet) (cytotoxin)	
	Haemophilus ducreyi	Cytolethal distending toxin (cdtA, cdtB, cdtC)	U53215
	Helicobacter pylori	Vacuolating toxin (vacA)	U07145
			U80067
		•	U80068
			AF077938
			AF077939
			AF077940
			AF077941
	Legionella programanhila	Structural toxin protein (rtxA)	AF057703
	Legionella pneumophila	Listeriolysin O (lisA, hlyA)	X15127
	Listeria monocytogenes	Listeriorysui O (iish, illyh)	M24199
			X60035
			U25452
			U25443
			U25446
			U25449
	Pasteurella multocida	Mitogenic toxin (dermonecrotic toxin)	X57775, Z2838
	i with the multitude	 	X51512
			X52478
	Destar misskille	Hemolysin (hpmA)	M30186
	Proteus mirabilis	Cytotoxin (Enterotoxin A)	X14956
	Pseudomonas aeruginosa Salmonella typhimurium	Calmodulin-sensitive adenylate cyalase toxin (cya)	
		Cytolysin (salmolysin) (slyA)	U03842
			L16014

Table 6. List of bacterial toxins selected for diagn stic purposes (continued).

	rganism	T xin	Accession number
	erratia marcescens	Hemolysin (shlA)	M22618
	iigella dysenteriae type 1	Shiga toxin (stxA and stxB)	X07903, M32511
			M19437
			M24352, M21947
C1	nigella flexneri	ShET2 enterotoxin (senA)	Z54211
Sh	ихена пелнен	Shell entitionin (som)	Z47381
		Enterotoxin 1 (set1A and set1B)	U35656
		Hemolysin E (hlyE, clyA, sheA)	AF200955
CI	ti alla samusi	Shiga toxin ($stxA$ and $stxB$)	AJ132761
	higella sonnei		L01270
	phingomonas paucimobilis	Beta-hemolysin (hlyA)	D42143
St	taphylococcus aureus	Gamma-hemolysin (hlg2)	
			L01055
		Enterotoxin	U93688
		Enterotoxin A (sea)	L22565, L22566
			M18970
		Enterotoxin B	M11118
		Enterotoxin C1 (entC1)	X05815
		Enterotoxin C2 (entC2)	P34071
		Enterotoxin C3 (entC3)	X51661
		Enterotoxin D (sed)	M94872
		Enterotoxin E	M21319
		Enterotoxin G (seg)	AF064773
		Enterotoxin H (seh)	U11702
		Enterotoxin I (sei)	AF064774
		Enterotoxin J	AF053140
		Exfoliative toxin A (ETA, Epidermolytic toxin A)	M17347
			M17357
			L25372, M20371
		Exfoliative toxin B (ETB)	M17348, M1377
		Leukocidin R (F and S component, lukF and lukS;	X64389, S53213
		Hemolysin B and C)	X72700
		Hemolysin B and C)	L01055
		m i i i i i i i i i i i i mnom i	V01645
		Toxic shock syndrome toxin 1 (TSST-1,	X01645
		alpha toxin, alpha hemolysin)	M90536
			J02615
			U93688
	Staphylococcus epidermidis	Delta toxin (hld)	AF068634
	Staphylococcus intermedius	Enterotoxin 1	U91526
1		Leukocidin R (F and S component, lukF and lukS;	X79188
		synergohymenotropic toxin)	
			X52474

Table 6. List f bacterial t xins selected f r diagn stic purposes (c ntinued).

Organism	Toxin	Accessi n number
Streptococcus pyogenes	Streptococcus pyrogenic exotoxin A (speA)	X61553 to X61573 X03929 U40453, M19350
	Pyrogenic exotoxin B (speB) M86905, M35110	U63134
Vibrio cholerae	Cholerae toxin (ctxA and ctxB subunits)	X00171 X76390 X58786 X58785, S55782 D30052 D30053 K02679 AF175708
	Accessory cholera enterotoxin (ace)	Z22569, AF1757
	Heat-stable enterotoxin (sto)	X74108, M85198 M97591, L03220
	Zonula occludens toxin (zot)	M83563, AF1757
Vibrio parahaemolyticus	Thermostable direct hemolysin (tdh)	S67841
Vibrio vulnificus	Cytolysin (vvhA)	M34670
Yersinia enterocolitica	Heat-stable enterotoxin (yst)	U09235, X65999
	Heat-stable enterotoxin type B (ystB)	D88145
	Heat-stable enterotoxin type C (ystC)	D63578
Yersinia kristensenii	Enterotoxin X69218	
Yersinia pestis	Toxin	X92727

Table 7. Origin f the nucleic acids and/ r sequences in the sequence listing.

	SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source Gene*	
_	1	Acinetobacter baumannii	This patent	tuf
	2	Actinomyces meyeri	This patent	tuf
	3	Aerococcus viridans	This patent	tuf
	4	Achromobacter xylosoxidans subsp. denitrificans	This patent	tuf
	5	Anaerorhabdus furcosus	This patent	tuf
)	6	Bacillus anthracis	This patent	tuf
	7	Bacillus cereus	This patent	tuf
	8	Bacteroides distasonis	This patent	tuf +f
	9	Enterococcus casseliflavus	This patent	tuf tuf
	10	Staphylococcus saprophyticus	This patent	tuf tuf
;	11	Bacteroides ovatus	This patent This patent	tuf
	12	Bartonella henselae	This patent	tuf
	13	Bifidobacterium adolescentis	This patent	tuf
	14	Bifidobacterium dentium	This patent	tuf
	15	Brucella abortus	This patent	tuf
)	16	Burkholderia cepacia Cedecea davisae	This patent	tuf
	17 18	Cedecea neteri	This patent	túf
	18	Cedecea lapagei	This patent	tuf
	20	Chlamydia pneumoniae	This patent	tuf
5	20 21	Chlamydia psittaci	This patent	tuf
,	22	Chlamydia trachomatis	This patent	tuf
	23	Chryseobacterium meningosepticum	This patent	tuf
	24	Citrobacter amalonaticus	This patent	tuf
	25	Citrobacter braakii	This patent	tuf
0	26	Citrobacter koseri	This patent	tuf
•	27	Citrobacter farmeri	This patent	tuf
	28	Citrobacter freundii	This patent	tuf
	29	Citrobacter sedlakii	This patent	tuf tuf
	30	Citrobacter werkmanii	This patent	tuf tuf
5	31	Citrobacter youngae	This patent This patent	tuf
	32	Clostridium perfringens	This patent	tuf
	33	Comamonas acidovorans	This patent	tuf
	34	Corynebacterium bovis	This patent	tuf
^	35	Corynebacterium cervicis	This patent	tuf
0	36	Corynebacterium flavescens Corynebacterium kutscheri	This patent	tuf
	37	Corynebacterium minutissimum	This patent	tuf
	38 39	Corynebacterium mycetoides	This patent	tuf
	40	Corynebacterium pseudogenitalium	This patent	tuf
5	41	Corynebacterium renale	This patent	tuf
	42	Corynebacterium ulcerans	This patent	tuf
	43	Corynebacterium urealyticum	This patent	tuf
	44	Corynebacterium xerosis	This patent	tuf
	45	Coxiella burnetii	This patent	tuf
50	46	Edwardsiella hoshinae	This patent	tuf tuf
	47	Edwardsiella tarda	This patent	tuf tuf
	48	Eikenella corrodens	This patent	tuf
	49	Enterobacter aerogenes	This patent This patent	tuf
	50	Enterobacter agglomerans	This patent	tuf
55	51	Enterobacter amnigenus	This patent	tuf
	52	Enterobacter asburiae	This patent	tuf
	53	Enterobacter cancerogenus	This patent	tuf
	54 55	Enterobacter cloacae	This patent	tuf
60	55	Enterobacter gergoviae Enterobacter hormaechei	This patent	túf
60	56 57	Enterobacter normaetnet Enterobacter sakazakii	This patent	tuf
	57 58	Enterococcus casseliflavus	This patent	tif
	58 59	Enterococcus cecorum	This patent	tuf
	60	Enterococcus dispar	This patent	tuf
65	61	Enterococcus durans	This patent	tuf

Table 7. Origin of the nucleic acids and/or sequences in the sequence listing (c ntinued).

•	SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
-	62	Enterococcus faecalis	This patent	tuf
	63	Enterococcus faecalis	This patent	tuf
	64	Enterococcus faecium	This patent	tuf
	65	Enterococcus flavescens	This patent	tuf
	66	Enterococcus gallinarum	This patent	tuf
	67	Enterococcus hirae	This patent	tuf
	68	Enterococcus mundtii	This patent	tuf
	69	Enterococcus pseudoavium	This patent	tuf
	70	Enterococcus raffinosus	This patent	tuf
	71	Enterococcus saccharolyticus	This patent	tuf tuf
	72	Enterococcus solitarius	This patent	tuf
	73	Enterococcus casseliflavus	This patent	tuf (C) unknow:
	74	Staphylococcus saprophyticus	This patent	tuf (C)
	75	Enterococcus flavescens	This patent	tuf (C)
	76	Enterococcus gallinarum	This patent This patent	tuf (C)
	· 77	Ehrlichia canis	This patent	tuf
	78	Escherichia coli	This patent	tuf
	79	Escherichia fergusonii	This patent	tuf
	80	Escherichia hermannii	This patent	tuf
	81	Escherichia vulneris	This patent	tuf
	82	Eubacterium lentum Eubacterium nodatum	This patent	tuf
	83	Ewingella americana	This patent	tuf
	84	Francisella tularensis	This patent	tuf
	85 86	Fusobacterium nucleatum subsp. polymorphum	This patent	tuf
		Gemella haemolysans	This patent	tuf
	87 88	Gemella morbillorum	This patent	tuf
	89	Haemophilus actinomycetemcomitans	This patent	tuf
	90	Haemophilus aphrophilus	This patent	tuf
	91	Haemophilus ducreyi	This patent	tuf
5	92	Haemophilus haemolyticus	This patent	tuf
•	93	Haemophilus parahaemolyticus	This patent	tuf
	94	Haemophilus parainfluenzae	This patent	tuf
	95	Haemophilus paraphrophilus	This patent	tuf
	96	Haemophilus segnis	This patent	tuf
)	97	Hafnia alvei	This patent	tuf
	98	Kingella kingae	This patent	tuf
	99	Klebsiella ornithinolytica	This patent	tuf ****
	100	Klebsiella oxytoca	This patent	tuf tuf
	101	Klebsiella planticola	This patent	tuf tuf
5	102	Klebsiella pneumoniae subsp. ozaenae	This patent This patent	tuf
	103	Klebsiella pneumoniae pneumoniae	This patent	tuf
	104	Klebsiella pneumoniae subsp. rhinoscleromatis	This patent	tuf
	105	Kluyvera ascorbata	This patent	tuf
_	106	Kluyvera cryocrescens	This patent	tuf
0	107	Kluyvera georgiana Lactobacillus casei subsp. casei	This patent	tuf
	108	Lactococcus lactis subsp. lactis	This patent	tuf
	109	Leclercia adecarboxylata	This patent	tuf
	110 111	Legionella micdadei	This patent	tuf
5	112	Legionella pneumophila subsp. pneumophila	This patent	tuf
,	113	Leminorella grimontii	This patent	tuf
	113	Leminorella richardii	This patent	tuf
	115	Leptospira interrogans	This patent	tuf
	116	Megamonas hypermegale	This patent	tuf
0	117	Mitsuokella multacidus	This patent	tuf
J	118	Mobiluncus curtisii subsp. holmesii	This patent	tuf
	119	Moellerella wisconsensis	This patent	tuf
	120	Moraxella catarrhalis	This patent	tuf mf
	121	Morganella morganii subsp. morganii	This patent	tuf
55	122	Mycobacterium tuberculosis	This patent	tuf

Table 7. Origin f the nucleic acids and/or sequences in the sequence listing (continued).

	SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source Gene*	
	123	Neisseria cinerea	This patent	tuf
	124	Neisseria elongata subsp. elongata	This patent	tuf
	125	Neisseria flavescens	This patent	tuf
	126	Neisseria gonorrhoeae	This patent	tuf
	127	Neisseria lactamica	This patent	tuf
	128	Neisseria meningitidis	This patent	tuf
	129	Neisseria mucosa	This patent	tuf
	130	Neisseria sicca	This patent	tuf
	131	Neisseria subflava	This patent	tuf
	132	Neisseria weaveri	This patent	tuf
	132	Ochrobactrum anthropi	This patent	tuf
	134	• • • • • • • • • • • • • • • • • • •	This patent	tuf
	135	Pantoea agglomerans Pantoea dispersa	This patent	tuf
	136	Pasteurella multocida	This patent	tuf
	130		This patent	tuf
		Peptostreptococcus anaerobius	This patent	tuf
	138	Peptostreptococcus asaccharolyticus		tuf
	139	Peptostreptococcus prevotii	This patent	
	140	Porphyromonas asaccharolytica	This patent	tuf tuf
	141	Porphyromonas gingivalis	This patent	tuf tuf
	142	Pragia fontium	This patent	tuf tuf
	143	Prevotella melaninogenica	This patent	tuf tuf
	144	Prevotella oralis	This patent	tuf
	145	Propionibacterium acnes	This patent	tuf
	146	Proteus mirabilis	This patent	tuf
	147	Proteus penneri	This patent	tuf
	148	Proteus vulgaris	This patent	tuf
	149	Providencia alcalifaciens	This patent	tuf
	150	Providencia rettgeri	This patent	tuf
	151	Providencia rustigianii	This patent	tuf
	152	Providencia stuartii	This patent	tuf
	153	Pseudomonas aeruginosa	This patent	tuf
	154	Pseudomonas fluorescens	This patent	tuf
	155	Pseudomonas stutzeri	This patent	tuf
	156	Psychrobacter phenylpyruvicum	This patent	tuf
	157	Rahnella aquatilis	This patent	tuf
)	. 158	Salmonella choleraesuis subsp.arizonae	This patent	tuf
	159	Salmonella choleraesuis subsp. choleraesuis	This patent	tuf
		serotype Choleraesuis	•	
	160	Salmonella choleraesuis subsp. diarizonae	This patent	tuf
	161	Salmonella choleraesuis subsp. choleraesuis	This patent	tuf
;		serotype Heidelberg	•	•
	162	Salmonella choleraesuis subsp. houtenae	This patent	tuf
	163	Salmonella choleraesuis subsp. indica	This patent	tuf
	164	Salmonella choleraesuis subsp. salamae	This patent	tuf
	165	Salmonella choleraesuis subsp. choleraesuis seroty		
)	166	Serratia fonticola	This patent	tuf
•	167	Serratia liquefaciens	This patent	tuf
	168	Serratia marcescens	This patent	tuf
	169	Serratia marcescens Serratia odorifera	This patent	tuf
	170	Serratia odorijera Serratia plymuthica	This patent	tuf
5		Serratia ptymumica Serratia rubidaea	This patent	tuf
,	171		This patent	tuf
	172	Shigella boydii Shigella disenteriae	This patent	tuf
	173	Shigella dysenteriae		
	174	Shigella flexneri	This patent	tuf tuf
`	175	Shigella sonnei	This patent	tuf tuf
)	176	Staphylococcus aureus	This patent	tuf
	177	Staphylococcus aureus	This patent	tuf
	178	Staphylococcus aureus	This patent	tuf
	179	Staphylococcus aureus	This patent	tuf
_	180	Staphylococcus aureus subsp. aureus	This patent	tuf
5	181	Staphylococcus auricularis	This patent	tuf
	182	Staphylococcus capitis subsp. capitis	This patent	tuf

Table 7. Origin of the nucleic acids and/or sequences in the sequence listing (continued).

	SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
_	183	Macrococcus caseolyticus	This patent	tuf
	184	Staphylococcus cohnii subsp. cohnii	This patent	tuf
	185	Staphylococcus epidermidis	This patent	tuf
	186	Staphylococcus haemolyticus	This patent	tuf
	187	Staphylococcus warneri	This patent	tuf
	188	Staphylococcus haemolyticus	This patent	tuf
	189	Staphylococcus haemolyticus	This patent	tuf
	190	Staphylococcus haemolyticus	This patent	tuf
	190	Staphylococcus hominis subsp. hominis	This patent	tuf
	192	Staphylococcus warneri	This patent	tuf
	193	Staphylococcus hominis	This patent	tuf
	194	Staphylococcus hominis	This patent	tuf
	195	Staphylococcus hominis	This patent	tuf
	196	Staphylococcus hominis	This patent	tuf
	197	Staphylococcus lugdunensis	This patent	tuf
	198	Staphylococcus saprophyticus	This patent	tuf
	199	Staphylococcus saprophyticus	This patent	tuf
	200	Staphylococcus saprophyticus	This patent	tuf
	201	Staphylococcus sciuri subsp. sciuri	This patent	tuf
	202	Staphylococcus warneri	This patent	tuf
	202	Staphylococcus warneri	This patent	tuf
	203	Bifidobacterium longum	This patent	tuf
	205	Stenotrophomonas maltophilia	This patent	tuf
	206	Streptococcus acidominimus	This patent	tuf
	207	Streptococcus agalactiae	This patent	tuf
	208	Streptococcus agalactiae	This patent	tuf
	209	Streptococcus agalactiae	This patent	tuf
	210	Streptococcus agalactiae	This patent	tuf
	211	Streptococcus anginosus	This patent	tuf
	212	Streptococcus bovis	This patent	tuf
	213	Streptococcus anginosus	This patent	tuf
	214	Streptococcus cricetus	This patent	tuf
	215	Streptococcus cristatus	This patent-	tuf
	216	Streptococcus downei	This patent	tuf
	217	Streptococcus dysgalactiae	This patent	tuf
	218	Streptococcus equi subsp. equi	This patent	tuf
	219	Streptococcus ferus	This patent	tuf
	220	Streptococcus gordonii	This patent	tuf
	221	Streptococcus anginosus	This patent	tuf
	222	Streptococcus macacae	This patent	tuf
	223	Streptococcus gordonii	This patent	tuf
	224	Streptococcus mutans	This patent	tuf
	225	Streptococcus parasanguinis	This patent	tuf
	226	Streptococcus ratti	This patent	tuf
	227	Streptococcus sanguinis	This patent	tuf
	228	Streptococcus sobrinus	This patent	tuf
	229	Streptococcus suis	This patent	tuf
	230	Streptococcus uberis	This patent	tuf
	231	Streptococcus vestibularis	This patent	tuf
	232	Tatumella ptyseos	This patent	tuf
	233	Trabulsiella guamensis	This patent	tuf
	234	Veillonella parvula	This patent	tuf
	235	Yersinia enterocolitica	This patent	tuf
	236	Yersinia frederiksenii	This patent	tuf
	237	Yersinia intermedia	This patent	tuf
)	238	Yersinia pestis	This patent	tuf
,	239	Yersinia pseudotuberculosis	This patent	tuf
	240	Yersinia rohdei	This patent	tuf
	240 241	Yokenella regensburgei	This patent	tuf
	241	Achromobacter xylosoxidans subsp. denitrificans	This patent	atpL
5	242	Acinetobacter baumannii	This patent	atpL
	7.44.7	Acinetobacter lwoffii	This patent	atpL

Table 7. Origin of the nucleic acids and/or sequences in the sequence listing (continued).

	SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
	245	Staphylococcus saprophyticus	This patent	atpD
	245	Alcaligenes faecalis subsp. faecalis	This patent	atpD
	246	Bacillus anthracis	This patent	atpD
	247	Bacillus cereus	This patent	atpD
	248	Bacteroides distasonis	This patent	atpD
	249		This patent	atpD
	250	Bacteroides ovatus	This patent	atpD
	251	Leclercia adecarboxylata	This patent	atpD
	252	Stenotrophomonas maltophilia	This patent	atpD
	253	Bartonella henselae	This patent	atpD
	254	Bifidobacterium adolescentis	This patent	atpD
	255	Brucella abortus	This patent	atpD
	256	Cedecea davisae	This patent	atpD
	257	Cedecea lapagei		atpD
	258	Cedecea neteri	This patent	atpD
	259	Chryseobacterium meningosepticum	This patent	atpD atpD
)	260	Citrobacter amalonaticus	This patent	
	261	Citrobacter braakii	This patent	atpD atpD
	262	Citrobacter koseri	This patent	
	263	Citrobacter farmeri	This patent	atpD
	264	Citrobacter freundii	This patent	atpD
,	265	Citrobacter koseri	This patent	atpD
	266	Citrobacter sedlakii	This patent	atpD
	267	Citrobacter werkmanii	This patent	atpD
	268	Citrobacter youngae	This patent	atpD
	269	Clostridium innocuum	This patent	atpD
)	270	Clostridium perfringens	This patent	atpD
	272	Corynebacterium diphtheriae	This patent	atpD_
	273	Corynebacterium pseudodiphtheriticum	This patent	atpD
	274	Corynebacterium ulcerans	This patent	atpD
	275	Corynebacterium urealyticum	This patent	atpD
5	276	Coxiella burnetii	This patent	atpD
,	277	Edwardsiella hoshinae	This patent	atpD
	278	Edwardsiella tarda	This patent	atpD
	279 279	Eikenella corrodens	This patent	atpD
	280	Enterobacter agglomerans	This patent	atpD
0	281	Enterobacter amnigenus	This patent	atpD
U	282	Enterobacter asburiae	This patent	atpD
		Enterobacter cancerogenus	This patent	atpD
	283	Enterobacter claacae	This patent	atpD
	284	Enterobacter gergoviae	This patent	atpD
_	285	Enterobacter gergovide Enterobacter hormaechei	This patent	atpD
5	286	Enterobacter normaechei Enterobacter sakazakii	This patent	atpD
	287		This patent	atpD
	288	Enterococcus avium	This patent	atpD
	289	Enterococcus casseliflavus	This patent	atpD
_	290	Enterococcus durans	This patent	atpD
0	291	Enterococcus faecalis	This patent	atpD
	292	Enterococcus faecium	This patent	atpD
	293	Enterococcus gallinarum	This patent	atpD
	294	Enterococcus saccharolyticus		atpD
_	295	Escherichia fergusonii	This patent	
5	296	Escherichia hermannii	This patent	atpD
	297	Escherichia vulneris	This patent	atpD
	298	Eubacterium lentum	This patent	atpD atpD
	299	Ewingella americana	This patent	atpD
	300	Francisella tularensis	This patent	atpD
0	301	Fusobacterium gonidiaformans	This patent	atpD
•	302	Fusobacterium necrophorum subsp. necrophorum	This patent	atpD
	303	Fusobacterium nucleatum subsp. polymorphum	This patent	atpD
	304	Gardnerella vaginalis	This patent	atpD
	305	Gemella haemolysans	This patent	atpD
55		Gemella morbillorum	This patent	atpD

Table 7. Origin of the nucleic acids and/ r sequences in the sequence listing (c ntinued).

S	EQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
	307	Haemophilus ducreyi	This patent	atpD
	308	Haemophilus haemolyticus	This patent	atpD
			This patent	atpD
	309	Haemophilus parahaemolyticus		atpD
	310	Haemophilus parainfluenzae	This patent	
	311	Hafnia alvei	This patent	atpD
	312	Kingella kingae	This patent	atpD
	313	Klebsiella pneumoniae subsp. ozaenae	This patent	atpD
	314	Klebsiella ornithinolytica	This patent	atpD
	315	Klebsiella oxytoca	This patent	atpD
	316	Klebsiella planticola	This patent	atpD
	317	Klebsiella pneumoniae subsp. pneumoniae	This patent	atpD
	318	Kluyvera ascorbata	This patent	atpD
	319	Kluyvera cryocrescens	This patent	atpD
	320	Kluyvera georgiana	This patent	atpD
	321	Lactobacillus acidophilus	This patent	atpD
	322	Legionella pneumophila subsp. pneumophila	This patent	atpD
	323	Leminorella grimontii	This patent	atpD
	324	Listeria monocytogenes	This patent	atpD
	325	Micrococcus lylae	This patent	atpD
	326	Moellerella wisconsensis	This patent	atpD
	327	Moraxella catarrhalis	This patent	atpD
		Moraxella osloensis	This patent	atpD
	328	*	This patent	atpD
	329	Morganella morganii subsp. morganii	This patent	atpD
	330	Pantoea agglomerans		atpD
	331	Pantoea dispersa	This patent	-
	332	Pasteurella multocida	This patent	atpD
	333	Pragia fontium	This patent	atpD
	334	Proteus mirabilis	This patent	atpD
	335	Proteus vulgaris	This patent	atpD
	336	Providencia alcalifaciens	This patent	atpD
	337	Providencia rettgeri	This patent	atpD
	338	Providencia rustigianii	This patent	atpD
	339	Providencia stuartii	This patent	atpD
	340	Psychrobacter phenylpyruvicum	This patent	atpD
	341	Rahnella aquatilis	This patent	atpD
	342	Salmonella choleraesuis subsp. arizonae	This patent	atpD
	343	Salmonella choleraesuis subsp. choleraesuis serotype Choleraesuis	This patent	atpD
	344	Salmonella choleraesuis subsp. diarizonae	This patent	atpD
	345	Salmonella choleraesuis subsp. houtenae	This patent	atpD
	346	Salmonella choleraesuis subsp. indica	This patent	atpD_
	347	Salmonella choleraesuis subsp. choleraesuis serotype Paratyphi A	This patent	atpD atpD
	348	Salmonella choleraesuis subsp. choleraesuis	This patent	шры
		serotype Paratyphi B	This petant	amD
	349	Salmonella choleraesuis subsp. salamae	This patent	atpD
	350	Salmonella choleraesuis subsp. choleraesuis serotype	Typhi This patent	atpD
	351	Salmonella choleraesuis subsp. choleraesuis serotype Typhimurium	This patent This patent	atpD atpD
	352	Salmonella choleraesuis subsp. choleraesuis serotype Virchow	This patent	atpD
	353	Serratia ficaria	This patent	atpD
	354	Serratia fonticola	This patent	atpD
	355	Serratia grimesii	•	
	356	Serratia liquefaciens	This patent	atpD
	357	Serratia marcescens	This patent	atpD
	358	Serratia odorifera	This patent	atpD
	359	Serratia plymuthica	This patent	atpD
	360	Serratia rubidaea	This patent	atpD
	361	Pseudomonas putida	This patent	atpD
	362	Shigella boydii	This patent	atpD
	363	Shigella dysenteriae	This patent	atpD

Table 7. Origin of the nucleic acids and/or sequences in the sequence listing (continued).

	SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source Gene*	·
_	364	Shigella flexneri	This patent	atpD
	365	Shigella sonnei	This patent	atpD
	366	Staphylococcus aureus	This patent	atpD
	367	Staphylococcus auricularis	This patent	atpD
	368	Staphylococcus capitis subsp. capitis	This patent	atpD
	369	Staphylococcus cohnii subsp. cohnii	This patent	atpD
	370	Staphylococcus epidermidis	This patent	atpD
	371	Staphylococcus haemolyticus	This patent	atpD
	372	Staphylococcus hominis subsp. hominis	This patent	atpD
	373	Staphylococcus hominis	This patent	atpD
	374	Staphylococcus lugdunensis	This patent	atpD
	375	Staphylococcus saprophyticus	This patent	atpD
	376	Staphylococcus simulans	This patent	atpD
	377	Staphylococcus warneri	This patent	atpD
	378	Streptococcus acidominimus	This patent	atpD
	379	Streptococcus agalactiae	This patent	atpD
	380	Streptococcus agalactiae	This patent	atpD
	381	Streptococcus agalactiae	This patent	atpD
	382	Streptococcus agalactiae	This patent	atpD
	383	Streptococcus agalactiae	This patent	atpD
	384	Streptococcus dysgalactiae	This patent	atpD
	385	Streptococcus equi subsp. equi	This patent	atpD
	386	Streptococcus anginosus	This patent	atpD
	38 7	Streptococcus salivarius	This patent	atpD
	388	Streptococcus suis	This patent	atpD
	389	Streptococcus uberis	This patent	atpD
	390	Tatumella ptyseos	This patent	atpD
	391	Trabulsiella guamensis	This patent	atpD
	392	Yersinia bercovieri	This patent	atpD
	393	Yersinia enterocolitica	This patent	atpD
	394	Yersinia frederiksenii	This patent	atpD
	395	Yersinia intermedia	This patent	atpD
	396	Yersinia pseudotuberculosis	This patent	atpD
	397	Yersinia rohdei	This patent	atpD
	398	Yokenella regensburgei	This patent	atpD
	399	Yarrowia lipolytica	This patent	tuf (EF-
	400	Absidia corymbifera	This patent	tuf (EF-
	401	Alternaria alternata	This patent	tuf (EF-
	402	Aspergillus flavus	This patent	tuf (EF-
	403	Aspergillus fumigatus	This patent	tuf (EF-
	404	Aspergillus fumigatus	This patent	tuf (EF-
	405	Aspergillus niger	This patent	tuf (EF-
	406	Blastoschizomyces capitatus	This patent	tuf (EF-
	407	Candida albicans	This patent	tuf (EF-
	408	Candida albicans	This patent	tuf (EF-
	409	Candida albicans	This patent	tuf (EF-
	410	Candida albicans	This patent	tuf (EF-
	411	Candida albicans	This patent	tuf (EF-
	412	Candida dubliniensis	This patent	tuf (EF-
	413	Candida catenulata	This patent	tuf (EF
	414	Candida dubliniensis	This patent	tuf (EF
	415	Candida dubliniensis	This patent	tuf (EF
	416	Candida famata	This patent	tuf (EF
	417	Candida glabrata	WO98/20157	tuf (EF
	417	Candida guilliermondii	This patent	tuf (EF
)	419	Candida haemulonii	This patent	tuf (EF
•	420	Candida inconspicua	This patent	tuf (EF
	421	Candida kefyr	This patent	tuf (EF
	421	Candida krusei	WO98/20157	tuf (EF
	423	Candida lambica	This patent	tuf (EF
5	424	Candida lusitaniae	This patent	tuf (EF
	747		This patent	tuf (EF

Table 7. Origin of the nucleic acids and/or sequences in the sequence listing (c ntinued).

	SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
	426	Candida parapsilosis	WO98/20157	tuf (EF-1
	427	Candida rugosa	This patent	tuf (EF-1
	428	Candida sphaerica	This patent	tuf (EF-1
	429	Candida tropicalis	WO98/20157	tuf (EF-1
	430	Candida utilis	This patent	tuf (EF-
	431	Candida viswanathii	This patent	tuf (EF-
	432	Candida zeylanoides	This patent	tuf (EF-
	433	Coccidioides immitis	This patent	tuf (EF-
	434	Cryptococcus albidus	This patent	tuf (EF- tuf (EF-
	435	Exophiala jeanselmei	This patent	tuf (EF-
	436	Fusarium oxysporum	This patent	tuf (EF-
	437	Geotrichum sp.	This patent	tuf (EF-
	438	Histoplasma capsulatum	This patent This patent	tuf (EF-
	439	Issatchenkia orientalis Kudrjanzev	This patent	tuf (EF-
	440	Malassezia furfur	This patent	tuf (EF-
	441	Malassezia pachydermatis	This patent	tuf (EF-
	442	Malbranchea filamentosa	This patent	tuf (EF-
	443	Metschnikowia pulcherrima Paecilomyces lilacinus	This patent	tuf (EF-
	444	Paracoccidioides brasiliensis	This patent	tuf (EF
	445 446	Penicillium marneffei	This patent	tuf (EF
	440 447	Pichia anomala	This patent	tuf (EF
	448	Pichia anomala	This patent	tuf (EF
	449	Pseudallescheria boydii	This patent	tuf (EF
	450	Rhizopus oryzae	This patent	tuf (EF
)	451	Rhodotorula minuta	This patent	tuf (EF
	452	Sporobolomyces salmonicolor	This patent	tuf (EF
	453	Sporothrix schenckii	This patent	tuf (EF
	454	Ŝtephanoascus ciferrii	This patent	tuf (EF
	455	Trichophyton mentagrophytes	This patent	tuf (EF
,	456	Trichosporon cutaneum	This patent	tuf (EF
	457	Wangiella dermatitidis	This patent	tuf (EF
	458	Aspergillus fumigatus	This patent	atpD atpD
	459	Blastoschizomyces capitatus	This patent	atpD
_	460	Candida albicans	This patent This patent	atpD
)	461	Candida dubliniensis	This patent	atpD
	462	Candida famata	This patent	atpD
	463	Candida glabrata	This patent	atpD
	464	Candida guilliermondii	This patent	atpD
5	465	Candida haemulonii Candida inconspicua	This patent	atpD
,	466 467	Candida kefyr	This patent	atpD
	468	Candida krusei	This patent	atpD
	469	Candida lambica	This patent	atpD
	470	Candida lusitaniae	This patent	atpD
0	471	Candida norvegensis	This patent	atpD
U	472	Candida parapsilosis	This patent	atpD
	473	Candida rugosa	This patent	atpD
	474	Candida sphaerica	This patent	atpD
	475	Candida tropicalis	This patent	atpD
5	476	Candida utilis	This patent	atpD
	477	Candida viswanathii	This patent	atpD atpD
	478	Candida zeylanoides	This patent	
	479	Coccidioides immitis	This patent	atpD atpD
_	480	Cryptococcus albidus	This patent	atpD atpD
0	481	Fusarium oxysporum	This patent	atpD atpD
	482	Geotrichum sp.	This patent	atpD
	483	Histoplasma capsulatum	This patent This patent	atpD
	484	Malassezia furfur	This patent	atpD
٠,-	485	Malassezia pachydermatis	This patent	atpD
55	486	Metschnikowia pulcherrima	This patent	atpD
	487	Penicillium marneffei	Parent	_F =

Table 7. Origin f the nucleic acids and/ r sequences in the sequence listing (c ntinued).

Section		SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
489 Pichia anomala 490 Rhodotorula minuta 491 Rhodotorula minuta 491 Rhodotorula minuta 492 Sporobolomyees salmonicolor This patent app 493 Sporobolomyees salmonicolor This patent app 494 Stephanoascus ciferrii 495 Trichoplyton mentagrophytes 496 Wangiella dematitidis This patent app 497 Yarrowia lipolytica 151 498 Aspergillus famigatus 499 Blastoschizomyees capitatus 500 Candida rugosa 500 Candida rugosa 501 Coccidiodes immitis 502 Fuscarium asysportum 503 Histoplasma capsulatum 504 Paracoccidiodes brasiliensis This patent auf (M) 505 Penicillium marneffei 506 Penicillium marneffei 507 Trichophyton mentagrophytes 508 Penicillium marneffei 509 Penicillium marneffei 500 Candida rugos 500 Penicillium marneffei 501 Trichophyton mentagrophytes 503 Penicillium marneffei 504 Paracoccidiodes brasiliensis This patent auf (M) 506 Penicillium marneffei 507 Trichophyton mentagrophytes 508 Penicillium marneffei 509 Beaten begenina 500 Penicillium marneffei 500 Penicillium marneffei 501 Trichophyton mentagrophytes 502 Trichophyton mentagrophytes 503 Trichophyton mentagrophytes 504 Paracoccidione 506 Penicillium marneffei 507 Trichophyton mentagrophytes 508 Penicillium marneffei 509 Beaten begenina 500 Trichophyton mentagrophytes 500 Trichophyton mentagrophytes 501 Trichophyton mentagrophytes 502 Fusional This patent auf (M) 503 Beaten begenina 504 Paracoccidional 505 Trichophyton mentagrophytes 506 Trichophyton mentagrophytes 507 Trichophyton mentagrophytes 508 Penicillium marneffei 509 Beaten begenina 500 Trichophyton mentagrophytes 500 Trichophyton mentagrophytes 501 Trichophyton mentagrophytes 502 Trichophyton mentagrophytes 503 Trichophyton mentagrophytes 504 Penicillium marneffei 505 Trichophyton mentagrophytes 506 Trichophyton mentagrophytes 507 Trichophyton mentagrophytes 508 Trichophyton mentagrophytes 509 Beaten to the penicage trichophyton mentagrophytes 500 Trichophyton mentagrophytes 501 Trichophyton mentagrophytes 502 Trichophyton mentagrophytes 503 Trichophyton mentagrophytes 504 Leishmania denitopica 505 Tric	5	488	Pichia anomala	This patent	atpD
490 Rhodotorla minuta 491 Rhodotorla minuta 491 Rhodotorla minuta 492 Sporobolomyces salmonicolor 10 493 Sporobolomyces salmonicolor 10 493 Sporobolomyces salmonicolor 10 494 Stephanoascus ciferrii 495 Trichoplyton mentagrophytes 496 Wangiella dermatitidis 497 Varrowia lipolytica 115 498 Aspergillus famiganus 499 Blastoschizomyces capitatus 115 498 Aspergillus famiganus 116 499 Blastoschizomyces capitatus 117 500 Candida rugosa 118 500 Candida rugosa 118 500 Candida rugosa 119 500 Candida rugosa 119 500 Candida rugosa 119 500 Candida rugosa 119 500 Candida rugosa 110 500 Penicillum marneffei 110 500 Penicillum marneffei 111 500 Fenicillum marneffei 112 500 Fenicillum marneffei 113 500 Fenicillum marneffei 114 500 Fenicillum marneffei 115 500 Fenicillum marneffei 116 Leishmania derhiopica 117 Fenicum deriver 118 500 Fenicillum marneffei 119 Fenicum deriver 119 500 Fenicillum marneffei 110 Fenicum deriver 110 Fenicum deriver 110 Fenicum		489	Pichia anomala		
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614 Enterococcus malodoratus This patent tuf (C) 65 615 Enterococcus durans This patent tuf (C)		613		This natent	tuf
65 615 Enterococcus durans This patent tuf (C)					
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Table 7. Origin f the nucleic acids and/ r sequences in the sequence listing (continued).

SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
617	Enterococcus dispar	This patent	tuf (C)
618	Enterococcus avium	This patent	tuf (C)
619	Saccharomyces cerevisiae	Database	tuf (M)
621	Enterococcus faecium	This patent	tuf (C)
622	Saccharomyces cerevisiae	This patent	tuf (EF-1
623	Cryptococcus neoformans	This patent	tuf (EF-1
624	Candida albicans	WO98/20157	tuf (EF-1
662	Corynebacterium diphtheriae	WO98/20157	tuf
663	Candida catenulata	This patent	atpD
665	Saccharomyces cerevisiae	Database	tuf (EF-1
666	Saccharomyces cerevisiae	Database	atpD
667	Trypanosoma cruzi	This patent	atpD
668	Corynebacterium glutamicum	Database	tuf
669	Escherichia coli	Database	atpD
670	Helicobacter pylori	Database	atpD
671	Clostridium acetobutylicum	Database	atpD
672	Cytophaga lytica	Database	atpD
673	Ehrlichia risticii	This patent	atpD
674	Vibrio cholerae	This patent	atpD
675	Vibrio cholerae	This patent	tuf
676	Leishmania enriettii	This patent	atpD
677	Babesia microti	This patent	tuf (EF-1
678	Cryptococcus neoformans	This patent	atpD
679	Cryptococcus neoformans	This patent	atpD
680	Cunninghamella bertholletiae	This patent	atpD
684	Candida tropicalis	Database	atpD (V)
685	Enterococcus hirae	Database	atpD (V)
686	Chlamydia pneumoniae	Database	atpD (V)
687	Halobacterium salinarum	Database	atpD (V)
688	Homo sapiens	Database	atpD (V)
689	Plasmodium falciparum	Database	atpD (V)
690	Saccharomyces cerevisiae	Database	atpD (V)
691	Schizosaccharomyces pombe	Database	atpD(V)
. 692	Trypanosoma congolense	Database	atpD (V)
693	Thermus thermophilus	Database	atpD (V)
698	Escherichia coli	WO98/20157	tuf
709	Borrelia burgdorferi	Database	atpD (V)
710	Treponema pallidum	Database	atpD (V)
711	Chlamydia trachomatis	Genome project	atpD (V
712	Enterococcus faecalis	Genome project	atpD (V
713	Methanosarcina barkeri	Database	atpD (V)
714	Methanococcus jannaschii	Database	atpD (V)
715	Porphyromonas gingivalis	Genome project	atpD (V
716	Streptococcus pneumoniae	Genome project	atpD (V)
717	Burkholderia mallei	This patent	tuf
718	Burkholderia pseudomallei	This patent	tuf
719	Clostridium beijerinckii	This patent	tuf
720	Clostridium innocuum	This patent	tuf
72 1	Clostridium novyi	This patent	tuf
722	Clostridium septicum	This patent	tuf
723	Clostridium tertium	This patent	tuf
724	Clostridium tetani	This patent	tuf
725	Enterococcus malodoratus	This patent	tuf
726	Enterococcus sulfureus	This patent	tuf
727	Lactococcus garvieae	This patent	tuf
728	Mycoplasma pirum	This patent	tuf
729	Mycoplasma salivarium	This patent	tuf
730	Neisseria polysaccharea	This patent	tuf
731	Salmonella choleraesuis subsp. choleraesuis	This patent	tuf
	serotype Enteritidis		

Table 7. Origin of the nucleic acids and/ r sequences in the sequence listing (continued).

	SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
			This patent	tuf
	732	Salmonella choleraesuis subsp. choleraesuis serotype Gallinarum	_	
	733	Salmonella choleraesuis subsp. choleraesuis serotype Paratyphi B	This patent	tuf
)	734	Salmonella choleraesuis subsp. choleraesuis serotype Virchow	This patent	tuf
,	735	Serratia grimesii	This patent	tuf
	736	Clostridium difficile	This patent	tuf
	730 737	Burkholderia pseudomallei	This patent	atpD
	738	Clostridium bifermentans	This patent	atpD
;	739	Clostridium beijerinckii	This patent	atpD
	740	Clostridium difficile	This patent	atpD
	741	Clostridium ramosum	This patent	atpD
	742	Clostridium septicum	This patent	atpD
	743	Clostridium tertium	This patent	atpD
)	744	Comamonas acidovorans	This patent	atpD
	745	Klebsiella pneumoniae subsp. rhinoscleromatis	This patent	atpD
	746	Neisseria canis	This patent	atpD
	747	Neisseria cinerea	This patent	atpD atpD
	748	Neisseria cuniculi	This patent	atpD
5	749	Neisseria elongata subsp. elongata	This patent	atpD
	750	Neisseria flavescens	This patent This patent	atpD
	751	Neisseria gonorrhoeae	This patent	atpD
	752	Neisseria gonorrhoeae	This patent	atpD
_	753	Neisseria lactamica	This patent	atpD
)	754	Neisseria meningitidis	This patent	atpD
	755	Neisseria mucosa	This patent	atpD
	756	Neisseria subflava	This patent	atpD
	757	Neisseria weaveri	This patent	atpD
_	758 758	Neisseria animalis	This patent	atpD
5	759 760	Proteus penneri Salmonella choleraesuis subsp. choleraesuis	This patent	atpD
	261	serotype Enteritidis	This patent	atpD
	761 762	Yersinia pestis Burkholderia mallei	This patent	atpD
Λ	762 763	Clostridium sordellii	This patent	aipD
0	763 764	Clostridium novyi	This patent	atpD
	765	Clostridium botulinum	This patent	atpD
	765 766	Clostridium histolyticum	This patent	atpD
	767	Peptostreptococcus prevotii	This patent	atpD
5	768	Absidia corymbifera	This patent	atpD
9	769	Alternaria alternata	This patent	atpD
	7 7 0	Aspergillus flavus	This patent	atpD
	771	Mucor circinelloides	This patent	atpD
	772	Piedraia hortai	This patent	atpD
0	773	Pseudallescheria boydii	This patent	atpD
Ŭ	774	Rhizopus oryzae	This patent	atpD
	775	Scopulariopsis koningii	This patent	atpD
	776	Trichophyton mentagrophytes	This patent	atpD
	<i>777</i>	Trichophyton tonsurans	This patent	atpD atpD
5	778	Trichosporon cutaneum	This patent	
	779	Cladophialophora carrionii	This patent	tuf (EF tuf (EF
	780	Cunninghamella bertholletiae	This patent	tuf (EF
	781	Curvularia lunata	This patent	tuf (EF
	782	Fonsecaea pedrosoi	This patent	tuf (EF
50		Microsporum audouinii	This patent This patent	tuf (EF
	784	Mucor circinelloides	This patent	tuf (EF
	785	Phialophora verrucosa	This patent	tuf (EF
	786	Saksenaea vasiformis	This patent	tuf (EF
	787	Syncephalastrum racemosum	This patent	tuf (EF
65		Trichophyton tonsurans	This patent	tuf (EF
	789	Trichophyton mentagrophytes	Per	

Table 7. Origin of the nucleic acids and/or sequences in the sequence listing (c ntinued).

	SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
			This patent	tuf (EF-1)
5	790	Bipolaris hawaiiensis	This patent This patent	tuf (M)
	791	Aspergillus fumigatus	This patent	tuf (M)
	792	Trichophyton mentagrophytes	This patent	atpD (V)
	827	Clostridium novyi	This patent	atpD (V)
)	828	Clostridium difficile Clostridium septicum	This patent	atpD (V)
,	829 830	Clostridium botulinum	This patent	atpD (V)
	831	Clostridium perfringens	This patent	atpD (V)
	832	Clostridium tetani	This patent	atpD (V)
	833	Streptococcus pyogenes	Database	atpD (V)
5	834	Babesia bovis	This patent	atpD (V)
	835	Cryptosporidium parvum	This patent	atpD (V)
	836	Leishmania infantum	This patent	atpD (V)
	837	Leishmania major	This patent	atpD (V)
	838	Leishmania tarentolae	This patent	atpD (V)
0	839	Trypanosoma brucei	This patent	atpD (V)
	840	Trypanosoma cruzi	This patent	tuf (EF-1)
	841	Trypanosoma cruzi	This patent	tuf (EF-1) tuf (EF-1)
	842	Trypanosoma cruzi	This patent	tuf (M)
_	843	Babesia bovis	This patent This patent	tuf (M)
5	844	Leishmania aethiopica	This patent	tuf (M)
	845	Leishmania amazonensis	This patent	tuf (M)
	846	Leishmania donovani	This patent	tuf (M)
	847	Leishmania infantum Leishmania enriettii	This patent	tuf (M)
Λ	848	Leishmania gerbilli	This patent	tuf (M)
0	849 850	Leishmania major	This patent	tuf (M)
	851	Leishmania mexicana	This patent	tuf (M)
	852	Leishmania tarentolae	This patent	tuf (M)
	853	Trypanosoma cruzi	This patent	tuf (M)
55	854	Trypanosoma cruzi	This patent	tuf (M)
	855	Trypanosoma cruzi	This patent	tuf (M)
	856	Babesia bigemina	This patent	atpD
	857	Babesia bovis	This patent	atpD
	858	Babesia microti	This patent	atpD
10	859	Leishmania guyanensis	This patent	atpD
	860	Leishmania mexicana	This patent	atpD atpD
	861	Leishmania tropica	This patent	atpD
	862	Leishmania tropica	This patent Database	tuf
	863	Bordetella pertussis	Database	tuf (EF-
1 5	864	Trypanosoma brucei brucei	This patent	tuf (EF-
	865	Cryptosporidium parvum	This patent	atpD
	866	Staphylococcus saprophyticus	This patent	atpD
	867	Zoogloea ramigera Staphylococcus saprophyticus	This patent	tuf
50	868 869	Enterococcus casseliflavus	This patent	tuf
JU	870	Enterococcus casseliflavus	This patent	tuf
	871	Enterococcus flavescens	This patent	tuf
	872	Enterococcus gallinarum	This patent	tuf
	872 873	Enterococcus gallinarum	This patent	tuf
55	874	Staphylococcus haemolyticus	This patent	tuf
,,	875	Staphylococcus epidermidis	This patent	tuf
	876	Staphylococcus epidermidis	This patent	tuf
	877	Staphylococcus epidermidis	This patent	tuf
	878	Staphylococcus epidermidis	This patent	tuf
60	879	Enterococcus gallinarum	This patent	tuf
	880	Pseudomonas aeruginosa	This patent	tuf tuf
	881	Enterococcus casseliflavus	This patent	tuf tuf
	882	Enterococcus casseliflavus	This patent	tuf tuf
	883	Enterococcus faecalis	This patent	tuf
65		Enterococcus faecalis	This patent	tuf
	885	Enterococcus faecium	This patent	•49

Table 7. Origin f the nucleic acids and/or sequences in the sequence listing (c ntinued).

	SEQ	ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
	886		ccus faecium	This patent	tuf
	887		ramigera	This patent	tuf
	888		ccus faecalis	This patent	tuf
	889		us fumigatus	This patent	atpD
	890		un marneffei	This patent	atpD
	891		cyces lilacinus	This patent	atpD
	892		um marneffei	This patent	atpD
	893		ix schenckii	This patent	atpD
	894	Malhran	chea filamentosa	This patent	atpD
	895		vyces Iilacinus	This patent	atpD
	896	Aspergil		This patent	atpD
	897		lus fumigatus	This patent	tuf (EF-1)
	898		um marneffei	This patent	tuf (EF-1)
	899	Piedraia		This patent	tuf (EF-1)
	900		nortal nyces lilacinus	This patent	tuf (EF-1)
)	901	Paracoc	cidioides brasiliensis	This patent	tuf (EF-1)
,	902	Sporoth	ix schenckii	This patent	tuf (EF-1)
	903		um marneffei	This patent	tuf (EF-1)
	903		ria lunata	This patent	tuf (M)
	905		lus niger	This patent	tuf (M)
,	905		s hawaiiensis	This patent	tuf (M)
'	907		llus flavus	This patent	tuf (M)
	908	Alterna	ia alternata	This patent	tuf (M)
	909	Panicilli	ium marneffei	This patent	tuf (M)
	910		ium marneffei	This patent	tuf (M)
)	910	918	Escherichia coli	Database	recA
		929	Bacteroides fragilis	This patent	atpD (V)
		929	Bacteroides distasonis	This patent	atpD(V)
		930	Porphyromonas asaccharolytica	This patent	atpD(V)
		931	Listeria monocytogenes	This patent	tuf
5		932	Saccharomyces cerevisiae	Database	recA (Rad5)
,		940	Saccharomyces cerevisiae	Database	recA (Dmc1
		941	Cryptococcus humicolus	This patent	atpD
		942	Escherichia coli	This patent	atpD
		942	Escherichia coli	This patent	atpD
0		943 944	Escherichia coli	This patent	atpD
J			Escherichia coli	This patent	atpD
		945 946	Neisseria polysaccharea	This patent	atpD
		940	Neisseria sicca	This patent	atpD
		947 948	Streptococcus mitis	This patent	atpD
5		948 949	Streptococcus mitis	This patent	atpD
,		949 950	Streptococcus mitis	This patent	atpD
		950 951	Streptococcus oralis	This patent	atpD
		952	Streptococcus oraus Streptococcus pneumoniae	This patent	atpD
		952 953	Streptococcus pneumoniae	This patent	atpD
0		953 954	Streptococcus preumoniae	This patent	atpD
U			Streptococcus pneumoniae	This patent	atpD
		955 956	Babesia microti	This patent	atpD (V)
			Entamoeba histolytica	This patent	atpD (V)
		957 958	Fusobacterium nucleatum subsp. polymorphum	This patent	atpD (V)
5		959	Leishmania aethiopica	This patent	atpD (V)
ر			Leishmania tropica	This patent	atpD (V)
		960	Leishmania guyanensis	This patent	atpD (V)
		961 962	Leishmania donovani	This patent	atpD (V)
			Leishmania hertigi	This patent	atpD (V)
^		963		This patent	atpD (V)
0		964	Leishmania mexicana Leishmania tropica	This patent	atpD (V)
		965	Peptostreptococcus anaerobius	This patent	atpD (V)
		966	Bordetella pertussis	This patent	tuf
		967		This patent	tuf
55		968	Bordetella pertussis Enterococcus columbae	This patent	tuf
• •		969	EMENOLULUS COMMUNIC	F	-

Table 7. Origin f the nucleic acids and/or sequences in the sequence listing (continued).

	SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source Gene*	
	970	Enterococcus flavescens	This patent	tuf
	970 971	Streptococcus pneumoniae	This patent	tuf
	972	Escherichia coli	This patent	tuf
	973	Escherichia coli	This patent	tuf
	974	Escherichia coli	This patent	tuf
	975	Escherichia coli	This patent	tuf
	976	Mycobacterium avium	This patent	tuf
	977	Streptococcus pneumoniae	This patent	tuf
	978	Mycobacterium gordonae	This patent	tuf
	979	Streptococcus pneumoniae	This patent	tuf
;	980	Mycobacterium tuberculosis	This patent	tuf
	981	Staphylococcus warneri	This patent	tuf
	982	Streptococcus mitis	This patent	tuf
	983	Streptococcus mitis	This patent	tuf
	984	Streptococcus mitis	This patent	tuf
)	985	Streptococcus oralis	This patent	tuf
	986	Streptococcus pneumoniae	This patent	tuf
	987	Enterococcus hirae	This patent	tuf (C
	988	Enterococcus mundtii	This patent	tuf (C
	989	Enterococcus raffinosus	This patent	tuf (C
5	990	Bacillus anthracis	This patent	recA
	991	Prevotella melaninogenica	This patent	recA
	992	Enterococcus casseliflavus	This patent	tuf
	993	Streptococcus pyogenes	Database	speA
	1002	Streptococcus pyogenes	WO98/20157	tuf
)	1003	Bacillus cereus	This patent	recA
	1004	Streptococcus pneumoniae	This patent	pbp1
	1005	Streptococcus pneumoniae	This patent	pbp1
	1006	Streptococcus pneumoniae	This patent	pbp1 pbp1
	1007	Streptococcus pneumoniae	This patent	pbp1
5	1008	Streptococcus pneumoniae	This patent	pbp 1
	1009	Streptococcus pneumoniae	This patent	pbp1
	1010	Streptococcus pneumoniae	This patent This patent	pbp1
	1011	Streptococcus pneumoniae	This patent	pbp1
_	1012	Streptococcus pneumoniae	This patent	pbp l
0	1013	Streptococcus pneumoniae	This patent	pbp l
	1014	Streptococcus pneumoniae	This patent	pbpl
	1015	Streptococcus pneumoniae	This patent	pbp
	1016	Streptococcus pneumoniae	This patent	pbp
_	1017	Streptococcus pneumoniae	This patent	pbp.
5	1018	Streptococcus pneumoniae	This patent	pbp
	1019	Streptococcus pneumoniae	This patent	pbp
	1020	Streptococcus pneumoniae	This patent	pbp:
	1021	Streptococcus pneumoniae	This patent	pbp.
^	1022	Streptococcus pneumoniae Streptococcus pneumoniae	This patent	pbp.
0	1023	Streptococcus pneumoniae	This patent	pbp.
	1024	Streptococcus pneumoniae	This patent	pbp
	1025	Streptococcus pneumoniae	This patent	pbp
	1026	Streptococcus pneumoniae	This patent	pbp
5	1027	Streptococcus pneumoniae	This patent	pbp
)	1028 1029	Streptococcus pneumoniae	This patent	pbp
	1030	Streptococcus pneumoniae	This patent	pbp
	1030	Streptococcus pneumoniae	This patent	pbp
	1031	Streptococcus pneumoniae	This patent	pbp
50	1032	Streptococcus pneumoniae	This patent	pbp
JU	1033	Streptococcus pneumoniae	This patent	pbp
	1034	Streptococcus pneumoniae	This patent	pbp
	1035	Streptococcus pneumoniae	This patent	pbp
	1036	Streptococcus pneumoniae	This patent	pbp
55		Cit observe Lucius	=	

Table 7. Origin of the nucleic acids and/or sequences in the sequence listing (continued).

SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
1038	Streptococcus pneumoniae	This patent	pbp2x
1039	Streptococcus pneumoniae	This patent	pbp2x
1040	Streptococcus pneumoniae	This patent	pbp2x
1041	Streptococcus pneumoniae	This patent	pbp2x
1042	Streptococcus pneumoniae	This patent	pbp2x pbp2x
1043	Streptococcus pneumoniae	This patent	pbp2x pbp2x
1044	Streptococcus pneumoniae	This patent	pbp2x pbp2x
1045	Streptococcus pneumoniae	This patent	pbp2x pbp2x
1046	Streptococcus pneumoniae	This patent	pbp2x pbp2x
1047	Streptococcus pneumoniae	This patent	pbp2x pbp2x
1048	Streptococcus pneumoniae	This patent	
1049	Enterococcus faecium	This patent	pbp2x vanA
1050	Enterococcus gallinarum		
1051		This patent	vanA vanA
1052	Enterococcus faecium	This patent	
1053	Enterococcus faecium	This patent	vanA
	Enterococcus faecium	This patent	vanA
1054	Enterococcus faecalis	This patent	vanA
1055	Enterococcus gallinarum	This patent	vanA
1056	Enterococcus faecium	This patent	vanA
1057	Enterococcus flavescens	This patent	vanA
1058	Enterococcus gallinarum	This patent	vanC1
1059	Enterococcus gallinarum	This patent	vanC1
1060	Enterococcus casseliflavus	This patent	vanC2
1061	Enterococcus casseliflavus	This patent	vanC2
1062	Enterococcus casseliflavus	This patent	vanC2
1063	Enterococcus casseliflavus	This patent	vanC2
1064	Enterococcus flavescens	This patent	vanC3
1065	Enterococcus flavescens	This patent	vanC3
1066	Enterococcus flavescens	This patent	vanC3
1067	Enterococcus faecium	This patent	vanXY
1068	Enterococcus faecium	This patent	vanXY
1069	Enterococcus faecium	This patent	vanXY
1070	Enterococcus faecalis	This patent	vanXY
1071	Enterococcus gallinarum	This patent	vanXY
1072	Enterococcus faecium	This patent	vanXY
1073	Enterococcus flavescens	This patent	vanXY
1074	Enterococcus faecium	This patent	vanXY
1075	Enterococcus gallinarum	This patent	vanXY
1076	Escherichia coli	Database	stx_t
1077	Escherichia coli	Database	stx_2
1093	Staphylococcus saprophyticus	This patent	unknow
1117	Enterococcus faecium	Database	vanB
1138	Enterococcus gallinarum	Database	vanC1
1139	Enterococcus faecium	Database	vanA
1140	Enterococcus casseliflavus	Database	vanC2
1141	Enterococcus faecium	Database	van HA X
1169	Streptococcus pneumoniae	Database	pbp1a
1172	Streptococcus pneumoniae	Database	pbp2b
1173	Streptococcus pneumoniae	Database	pbp2x
1178	Staphylococcus aureus	Database	mecA
1183	Streptococcus pneumoniae	Database	hexA
1184	Streptococcus pneumoniae	This patent	hexA
1185	Streptococcus pneumoniae	This patent	hexA
1186	Streptococcus pneumoniae	This patent	hexA
1187	Streptococcus pneumoniae	This patent	hexA

Tabl 7. Origin of the nucl ic acids and/or sequences in the sequence listing (continued).

SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
1188	Streptococcus oralis	This patent	hexA
1189	Streptococcus mitis	This patent	hexA
1190	Streptococcus mitis	This patent	hexA
1191	Streptococcus mitis	This patent	hexA
1198	Staphylococcus saprophyticus	This patent	unknown
1215	Streptococcus pyogenes	Database	рср
1230	Escherichia coli	Database Database	tuf (EF-G)
1242	Enterococcus faecium	Database	ddl
1243	Enterococcus faecalis	Database	mtlF, mtlD
			unknown
1244	Staphylococcus aureus subsp. aureus	This patent	
1245	Bacillus anthracis	This patent	atpD
1246	Bacillus mycoides	This patent	atpD
1247	Bacillus thuringiensis	This patent	atpD
1248	Bacillus thuringiensis	This patent	atpD
1249	Bacillus thuringiensis	This patent	atpD
1250	Bacillus weihenstephanensis	This patent	atpD
1251	Bacillus thuringiensis	This patent	atpD
1252	Bacillus thuringiensis	This patent	atpD
1253	Bacillus cereus	This patent	atpD
1254	Bacillus cereus	This patent	atpD
1255	Staphylococcus aureus	This patent	gyrA
1256	Bacillus weihenstephanensis	This patent	atpD
1257	Bacillus anthracis	This patent	atpD
1258	Bacillus thuringiensis	This patent	atpD
1259	Bacillus cereus	This patent	atpD
1260	Bacillus cereus	This patent	atpD
1261	Bacillus thuringiensis	This patent	atpD
1262	Bacillus thuringiensis	This patent	atpD
1263	Bacillus thuringiensis	This patent	atpD
1264	Bacillus thuringiensis	This patent	atpD
1265	Bacillus anthracis	This patent	atpD
1266	Paracoccidioides brasiliensis	This patent	tuf (EF-1)
1267	Blastomyces dermatitidis	This patent	tuf (EF-1)
1268	Histoplasma capsulatum	This patent	tuf (EF-1)
1269	Trichophyton rubrum	This patent	tuf (EF-1)
1270	Microsporum canis	This patent	tuf (EF-1)
1271	Aspergillus versicolor	This patent	tuf (EF-1)
1272	Exophiala moniliae	This patent	tuf (EF-1)
1273	Hortaea wemeckii	This patent	tuf (EF-1)
1274	Fusarium solani	This patent	tuf (EF-1)
1275	Aureobasidium pullulans	This patent	tuf (EF-1)
1276	Blastomyces dermatitidis	This patent	tuf (EF-1)
1277	Exophiala dermatitidis	This patent	tuf (EF-1
1278	Fusarium moniliforme	This patent	tuf (EF-1
1279	Aspergillus terreus	This patent	tuf (EF-1
1280	Aspergillus fumigatus	This patent	tuf (EF-1
1281	Cryptococcus laurentii	This patent	tuf (EF-1
1282	Emmonsia parva	This patent	tuf (EF-1
1283	Fusarium solani	This patent	tuf (EF-1
1284	Sporothrix schenckii	This patent	tuf (EF-1
1285	Aspergillus nidulans	This patent	tuf (EF-1
1286	Cladophialophora carrionii	This patent	tuf (EF-1
1287	Exserohilum rostratum	This patent	tuf (EF-1
1288	Bacillus thuringiensis	This patent	recA
	_		recA
1289	Bacillus thuringiensis	This patent	
1299	Staphylococcus aureus	Databas	gyrA
1300	Escherichia coli	Database	gyrA
1307	Staphylococcus aureus	Database	gyrB
1320	Escherichia coli	Database	parC (grl
1321	Staphylococcus aureus	Database	parC (grl
1328	Staphylococcus aureus	Database	parE (grli

Table 7. Origin of th nucleic acids and/or s qu nces in th s qu nc listing (continued).

-	SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
_	1348	unidentified bacterium	Database	aac2la
	1351	Pseudomonas aeruginosa	Database	aac3lb
	1356	Serratia marcescens	Database	aac3llb
	1361	Escherichia coli	Database	aac3lVa
	1366	Enterobacter cloacae	Database	aac3Vla
	1371	Citrobacter koseri	Database	aac6la
	1376	Serratia marcescens	Database	aac6lc
		Escherichia coli	Database	ant3la
	1381	Staphylococcus aureus	Database	ant4la
	1386	Escherichia coli	Database	aph3la
	1391	Escherichia coli	Database	aph3lla
	1396	Enterococcus faecalis	Database	aph3IIIa
	1401	Acinetobacter baumannii	Database	aph3Vla
	1406		Database	blaCARB
	1411	Pseudomonas aeruginosa	Database	blaCMY-2
	1416	Klebsiella pneumoniae	Database	blaCTX-M-1
	1423	Escherichia coli Salmonella choleraesuis subsp. choleraesuis serotype	Database	blaCTX-M-2
	1428			
		Typhimurium	Database	blaIMP
	1433	Pseudomonas aeruginosa	Database	blaOXA2
	1438	Escherichia coli	Database	blaOXA10
	1439	Pseudomonas aeruginosa	Database	blaPER1
	1442	Pseudomonas aeruginosa	Database	blaPER2
	1445	Salmonella choleraesuis subsp. choleraesuis serotype	Daiabass	
		Typhimurium	Database	dfrA
	1452	Staphylococcus epidermidis	Database	dhfrla
	1461	Escherichia coli	Database	dhfrlb
	1470	Escherichia coli	Database	dhfrV
	1475	Escherichia coli	Database	dhfrVI
	1480	Proteus mirabilis	Database	dhfrVII
	1489	Escherichia coli	Database	dhfrVIII
	1494	Escherichia coli	Database	dhfrlX
	1499	Escherichia coli	Database	dhfrXII
	1504	Escherichia coli	Database	dhfrXIII
	1507	Escherichia coli	Database	dhfrXV
	1512	Escherichia coli	Database	dhfrXVII
	1517	Escherichia coli	This patent	fusA
	1518	Acinetobacter Iwoffii	This patent	fusA-tuf space
	1519	Acinetobacter lwoffii	This patent	tuf
	1520	Acinetobacter Iwoffii	This patent	fusA
	1521	Haemophilus influenzae	•	fusA-tuf space
	1522	Haemophilus influenzae	This patent	tuf
	1523	Haemophilus influenzae	This patent	fusA
	1524	Proteus mirabilis	This patent This patent	fusA-tuf space
	1525	Proteus mirabilis		tuf
	1526	Proteus mirabilis	This patent	atpD
	1527	Campylobacter curvus	This patent	өгеА
	1530	Escherichia coli	Database	ereB
	1535	Escherichia coli	Database	linA
	1540	Staphylococcus haemolyticus	Database	linB
	1545	Enterococcus faecium	Database	mefA
	1548	Streptococcus pyogenes	Database	
	1551	Streptococcus pneumoniae	Database	mefE
	1560	Escherichia coli	Database	mphA
	1561	Candida albicans	This patent	tuf (EF-1)
	1562	Candida dubliniensis	This patent	tuf (EF-1)
	1563	Candida famata	This patent	tuf (EF-1)
	1564	Candida glabrata	This patent	tuf (EF-1)
	1565	Candida guilliermondii	This patent	tuf (EF-1)
	1566	Candida haemulonii	This patent	tuf (EF-1)
	1567	Candida kefyr	This patent	tuf (EF-1)
		Candida lusitaniae	This patent	tuf (EF-1)

Tabl 7. Origin of the nucleic acids and/or s qu nc s in the sequ nce listing (continu d).

	SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
	1569	Candida sphaerica	This patent	tuf (EF-1)
	1570	Candida tropicalis	This patent	tuf (EF-1)
	1571	Candida viswanathii	This patent	tuf (EF-1)
	1572	Alcaligenes faecalis subsp. faecalis	This patent	tuf
	1573	Prevotella buccalis	This patent	tuf
	1574	Succinivibrio dextrinosolvens	This patent	tuf
	1575	Tetragenococcus halophilus	This patent	tuf
	1576	Campylobacter jejuni subsp. jejuni	This patent	atpD
	1577	Campylobacter rectus	This patent	atpD
	1578	Enterococcus casseliflavus	This patent	fusA
	1579	Enterococcus gallinarum	This patent	fusA
	1580	Streptococcus mitis	This patent	fusA
	1585	Enterococcus faecium	Database	satG
	1590	Cloning vector pFW16	Database	tetM
	1594	Enterococcus faecium	Database	<i>van</i> D
	1599	Enterococcus faecalis	Database	vanE
	1600	Campylobacter jejuni subsp. doylei	This patent	atpD
	1601	Enterococcus sulfureus	This patent	atpD
	1602	Enterococcus solitarius	This patent	atpD
	1603	Campylobacter sputorum subsp. sputorum	This patent	atpD
	1604	Enterococcus pseudoavium	This patent	atpD
	1607	Klebsiella omithinolytica	This patent	gyrA
	1608	Klebsiella oxytoca	This patent	gyrA
	1613	Staphylococcus aureus	Database	vatB
	1618	Staphylococcus cohnii	Database	vatC
	1623	Staphylococcus aureus	Database	vga
	1628	Staphylococcus aureus	Database	vgaB
	1633	Staphylococcus aureus	Database	vgb
	1638	Aspergillus fumigatus	This patent	atpD
	1639	Aspergillus fumigatus	This patent	atpD
	1640	Bacillus mycoides	This patent	atpD
	1641	Bacillus mycoides	This patent	atpD
	1642	Bacillus mycoides	This patent	atpD
	1643	Bacillus pseudomycoides	This patent	atpD
	1644	Bacillus pseudomycoides	This patent	atpD
	1645	Budvicia aquatica	This patent	atpD
	1646	Buttiauxella agrestis	This patent	atpD
	1647	Candida norvegica	This patent	atpD
	1648	Streptococcus pneumoniae	This patent	pbp1a
	1649	Campylobacter lari	This patent	atpD
	1650	Coccidioides immitis	This patent	atpD
	1651	Emmonsia parva	This patent	atpD
	1652	Erwinia amylovora	This patent	atpD atpD
	1653	Fonsecaea pedrosoi	This patent	atpD
	1654	Fusarium moniliforme	This patent	aφD atpD
	1655	Klebsiella oxytoca	This patent	atpD atpD
	1656	Microsporum audouinii	This patent	atpD
	1657	Obesumbacterium proteus	This patent	афD atpD
	1658	Paracoccidioides brasiliensis	This patent This patent	atpD atpD
	1659	Plesiomonas shigelloides	This patent	atpD
	1660	Shewanella putrefaciens	This patent	tuf
	1662	Campylobacter curvus	This patent	tuf
	1663	Campylobacter rectus		tuf
	1664	Fonsecaea pedrosoi	This patent	tuf
	1666	Microsporum audouinii	This patent This patent	tuf
)	1667	Piedraia hortai		tuf
	1668	Escherichia coli	Database This patent	tuf
	1669	Saksenaea vasiformis	This patent	tuf
	1670	Trichophyton tonsurans	This pat nt	atpD
	1671	Enterobacter aerogenes	This patent	αφυ atpD
5	1672	Bordetella pertussis	Databas This sat, st	*
	1673	Arcanobacterium haemolyticum	This pat int	tuf

Tabl 7. Origin f th nucl ic acids and/ rs quenc s in th sequ nc listing (c ntinu d).

SEQ ID N	D. Archaeal, bacterial, funga	l or parasitical species Source	Gene*
1674	Butyrivibrio fibrisolvens	This patent	tuf
1674	Campylobacter jejuni subsp.		tuf
1675	Campylobacter lari	This patent	tuf
1676	Campylobacter sputorum su		tuf
1677		This patent	tuf
1678	Campylobacter upsaliensis	This patent	tuf .
1679	Globicatella sanguis	This patent	tuf
1680	Lactobacillus acidophilus	· · · · · · · · · · · · · · · · · · ·	tuf
1681	Leuconostoc mesenteroides	This patent	tuf
1682	Prevotella buccalis	This patent	tuf
1683	Ruminococcus bromii		atpD
1684	Paracoccidioides brasilierisi	This patent	tuf (EF-1)
1685	Candida norvegica	This patent	tuf
1686	Aspergillus nidulans	This patent	tuf
1687	Aspergillus terreus	This patent	tuf
1688	Candida norvegica	This patent	tuf
1689	Candida parapsilosis	WO98/20157	
1702	Streptococcus gordonii	WO98/20157	
1703	Streptococcus mutans	14/000/004/57	
1704	Streptococcus pneumoniae	WO98/20157	
1705	Streptococcus pyogenes		
1706	Streptococcus salivarius su	WO98/20157 WO98/20157	
1707	Escherichia coli	WO98/20157	
1708	Enterococcus faecalis	WO98/20157 WO98/20157	
1709	Pseudomonas aeruginosa	WO98/2015	
1710	Staphylococcus aureus	WO98/20151 WO98/20157	_
171	Escherichia coli	WO98/2015	
171:	Staphylococcus aureus	WO98/2015	_
171:	Enterococcus faecalis		recA
171		. jejuni WO98/2015	<i>_</i>
171	Abiotrophia adiacens	WO98/2015	_
171		WO98/2015 WO98/2015	
171		W000/001E	
171		m WO30/2013 WO98/2015	
171	Corynebacterium jeikeium		
172	Corynebacterium pseudod	WO98/2015	
172		WO98/2015	_
172		WO98/2015 WO98/2015	
172		WO98/2015	
172		WO98/2015 WO98/2015	
172		WQ98/2015	
172		WO98/2015 WO98/2015	
172		WO98/2013 WO98/2015	_
172			_
172	Staphylococcus saprophyt	w098/2015 W098/2015	_
173		WO98/2015 WO98/2015	· _
173	Streptococcus agalactiae		
173		WO98/2015 WO98/2015	
173	Streptococcus salivarius		
173	Agrobacterium radiobacter	WO98/2015 WO98/2015	·
173		WO98/2015 WO98/2015	
173	Bacteroides fragilis	WO98/2015 WO98/2015	
173		WO98/2015	
173			
173		WO98/2015 WO98/2015	_
174) Fibrobacter succinogenes		
174			
174	2 Helicobacter pylori	WO98/2015 WO98/2015	
174	3 Micrococcus luteus		
174		w098/2015 W098/2015	
174		WO98/2015 WO98/2015	
17	6 Neisseria gonorrhoeae	44.03-0150 L	,

Table 7. Origin f th nucleic acids and/or s qu nces in the s quence listing (continued).

SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
4747	Diskattaia prowazakii	WO98/20157	tuf
1747	Rickettsia prowazekii Salmonella choleraesuis subsp. choleraesuis	WO98/20157	tuf
1748		11000/20:01	
	serotype Typhimurium	WO98/20157	tuf
1749	Shewanella putrefaciens	WO98/20157 WO98/20157	tuf
1750	Stigmatella aurantiaca		tuf
1751	Thiomonas cuprina	WO98/20157	
1752	Treponema pallidum	WO98/20157	tuf
1753	Ureaplasma urealyticum	WO98/20157	tuf
1754	Wolinella succinogenes	WO98/20157	tuf
1755	Burkholderia cepacia	WO98/20157	tuf
1756	Bacillus anthracis	This patent	recA
1757	Bacillus anthracis	This patent	recA
1758	Bacillus cereus	This patent	recA
1759	Bacillus cereus	This patent	recA
1760	Bacillus mycoides	This patent	recA
	Bacillus pseudomycoides	This patent	recA
1761		This patent	recA
1762	Bacillus thuringiensis	This patent	recA
1763	Bacillus thuringiensis		gyrA
1764	Klebsiella oxytoca	This patent	
1765	Klebsiella pneumoniae subsp. ozaenae	This patent	gyrA
1766	Klebsiella planticola	This patent	gyrA
1767	Klebsiella pneumoniae	This patent	gyrA
1768	Klebsiella pneumoniae subsp. pneumoniae	This patent	gyrA
1769	Klebsiella pneumoniae subsp. pneumoniae	This patent	gyrA
1770	Klebsiella pneumoniae subsp. rhinoscleromatis	This patent	gyrA
1771	Klebsiella terrigena	This patent	gyrA
1772	Legionella pneumophila subsp. pneumophila	This patent	gyrA
1772	Proteus mirabilis	This patent	gyrA
		This patent	gyrA
1774	Providencia rettgeri	This patent	gyrA
1775	Proteus vulgaris	-	gyrA
1776	Yersinia enterocolitica	This patent	parC (grlA)
1777	Klebsiella oxytoca	This patent	
1778	Klebsiella oxytoca	This patent	parC (grlA)
1779	Klebsiella pneumoniae subsp. ozaenae	This patent	parC (grlA)
1780	Klebsiella planticola	This patent	parC (grlA)
1781	Klebsiella pneumoniae	This patent	parC (grlA)
1782	Klebsiella pneumoniae subsp. pneumoniae	This patent	parC (grlA)
1783	Klebsiella pneumoniae subsp. pneumoniae	This patent	parC (grlA)
1784	Klebsiella pneumoniae subsp. rhinoscleromatis	This patent	parC (grlA)
	Klebsiella terrigena	This patent	parC (grlA)
1785		This patent	fusA `
1786	Bacillus cereus	This patent	fusA
1787	Bacillus cereus	This patent	fusA
1788	Bacillus anthracis		fusA
1789	Bacillus cereus	This patent	fusA
1790	Bacillus anthracis	This patent	
1791	Bacillus pseudomycoides	This patent	fusA
1792	Bacillus cereus	This patent	fusA
1793	Bacillus anthracis	This patent	fusA
1794	Bacillus cereus	This patent	fusA
1795	Bacillus weihenstephanensis	This patent	fusA
1796	Bacillus mycoides	This patent	fusA
1797	Bacillus thuringiensis	This patent	fusA
	Bacillus weihenstephanensis	This patent	fusA-tuf spa
1798	and the second s	This patent	fusA-tuf spa
1799	Bacillus thuringiensis	This patent	fusA-tuf spa
1800	Bacillus anthracis	This patent	fusA-tuf spa
1801	Bacillus pseudomycoides		fusA-tuf spa
1802	Bacillus anthracis	This patent	•
1803	Bacillus cereus	This patent	fusA-tuf spa
1804	Bacillus cereus	This patent	fusA-tuf spa
1805	Bacillus mycoides	This patent	fusA-tuf spa
	Bacillus cereus	This patent	fusA-tuf spa

Table 7. Origin of the nucl ic acids and/ rs quenc s in the sequ nc listing (continued).

	SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
	1807	Bacillus cereus	This patent	fusA-tuf space
	1808	Bacillus cereus	This patent	fusA-tuf spacer
	1809	Bacillus anthracis	This patent	fusA-tuf space
	1810	Bacillus mycoides	This patent	tuf
	1811	Bacillus thuringiensis	This patent	tuf
	1812	Bacillus cereus	This patent	tuf
	1813	Bacillus weihenstephanensis	This patent	tuf
	1814	Bacillus anthracis	This patent	tuf
	1815	Bacillus cereus	This patent	tuf
	1816	Bacillus cereus	This patent	tuf
	1817	Bacillus anthracis	This patent	tuf
	1818	Bacillus cereus	This patent	tuf
	1819	Bacillus anthracis	This patent	tuf
	1820	Bacillus pseudomycoides	This patent	tuf
	1821	Bacillus cereus	This patent	tuf
	1822	Streptococcus oralis	This patent	fusA
	1823	Budvicia aquatica	This patent	fusA
	1824	Buttiauxella agrestis	This patent	fusA
	1825	Klebsiella oxytoca	This patent	fusA
	1826	Plesiomonas shigelloides	This patent	fusA
	1827	Shewanella putrefaciens	This patent	fusA
	1828	Obesumbacterium proteus	This patent	fusA
	1829	Klebsiella oxytoca	This patent	fusA-tuf space
	1830	Budvicia aquatica	This patent	fusA-tuf space
	1831	Plesiomonas shigelloides	This patent	fusA-tuf space
	1832	Obesumbacterium proteus	This patent	fusA-tuf space
	1833	Shewanella putrefaciens	This patent	fusA-tuf space
	1834	Buttiauxella agrestis	This patent	fusA-tuf space
	1835	Campylobacter coli	This patent	tuf
	1836	Campylobacter fetus subsp. fetus	This patent	tuf
	1837	Campylobacter fetus subsp. venerealis	This patent	tuf
	1838	Buttiauxella agrestis	This patent	tuf
	1839	Klebsiella oxytoca	This patent	tuf
	1840	Plesiomonas shigelloides	This patent	tuf
	1841	Shewanella putrefaciens	This patent	tuf
	1842	Obesumbacterium proteus	This patent	tuf
•	1843	Budvicia aquatica	This patent	tuf
	1844	Abiotrophia adiacens	This patent	atpD
	1845	Arcanobacterium haemolyticum	This patent	atpD
	1846	Basidiobolus ranarum	This patent	atpD
	1847	Blastomyces dermatitidis	This patent	atpD
	1848	Blastomyces dermatitidis	This patent	atpD
	1849	Campylobacter coli	This patent	atpD
	1850	Campylobacter fetus subsp. fetus	This patent	atpD
	1851	Campylobacter fetus subsp. venerealis	This patent	atpD
	1852	Campylobacter gracilis	This patent	atpD
		Campylobacter jejuni subsp. jejuni	This patent	atpD
	1853	Enterococcus cecorum	This patent	atpD
	1854		This patent	atpD
	1855	Enterococcus columbae	This patent	atpD
	1856	Enterococcus dispar	This patent	atpD
	1857	Enterococcus malodoratus	This patent	atpD atpD
	1858	Enterococcus mundtii	This patent	atpD
	1859	Enterococcus raffinosus		atpD atpD
	1860	Globicatella sanguis	This patent	
	1861	Lactococcus garvieae	This patent	atpD
	1862	Lactococcus lactis	This patent	atpD
	1863	Listeria ivanovii	This patent	atpD
	1864	Succinivibrio dextrinosolvens	This patent	atpD
	1865	Tetragenococcus halophilus	This patent	atpD
	1866	Campylobacter fetus subsp. fetus	This patent	recA
	1867	Campylobacter fetus subsp. venerealis	This patent	recA
	1868	Campylobacter jejuni subsp. jejuni	This patent	recA

Tabl 7. Origin of the nucl ic acids and/or sequenc s in the sequence listing (continued).

SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*	
1000	Enterococcus avium	This patent	recA	
1869	Enterococcus faecium	This patent	recA	
1870		This patent	recA	
1871	Listeria monocytogenes	This patent	recA	
1872	Streptococcus mitis	This patent	recA	
1873	Streptococcus oralis	This patent	tuf (M)	
1874	Aspergillus fumigatus	This patent	tuf (M)	
1875	Aspergillus versicolor	This patent	tuf (M)	
1876	Basidiobolus ranarum Campylobacter gracilis	This patent	tuf	
1877	Campylobacter jejuni subsp. jejuni	This patent	tuf	
1878	Coccidioides immitis	This patent	tuf (M)	
1879		This patent	tuf` ´	
1880	Erwinia amylovora Salmonella choleraesuis subsp. choleraesuis serotype	This patent	tuf	
1881		Tille parent	-	
4000	Typhimurium	Database	blaSHV	
1899	Klebsiella pneumoniae	Database	blaSHV	
1900	Klebsiella pneumoniae Escharishia soli	Database	blaSHV	
1901	Escherichia coli Klabriolla ppeumoniae	Database	blaSHV	
1902	Klebsiella pneumoniae Klebsiella pneumoniae	Database	blaSHV	
1903	Klebsiella pneumoniae Escherichia coli	Database	blaSHV	
1904	Pseudomonas aeruginosa	Database	blaSHV	
1905	Neisseria meningitidis	Database	blaTEM	
1927	Escherichia coli	Database	blaTEM	
1928		Database	blaTEM	
1929	Klebsiella oxytoca Escherichia coli	Database	blaTEM	
1930	Escherichia coli	Database	blaTEM	
1931	Escherichia coli	Database	blaTEM	
1932	Escherichia coli	Database	blaTEM	
1933 1954	Klebsiella pneumoniae subsp. pneumoniae	Database	gyrA	
	Candida inconspicua	This patent	tuf (M)	
1956 1957	Candida utilis	This patent	tuf (M)	
	Candida unis Candida zeylanoides	This patent	tuf (M)	
1958 1959	Candida zayianoides Candida catenulata	This patent	tuf (M)	
1960	Candida krusei	This patent	tuf (M)	
1965	Plasmid pGS05	Database	sulli	
1970	Transposon Tn10	Database	tetB	
1985	Cryptococcus neoformans	Database	tuf (EF-1	
1986	Cryptococcus neoformans	Database	tuf (EF-1	
1987	Saccharomyces cerevisiae	Database	tuf (EF-1	
1988	Saccharomyces cerevisiae	Database	tuf (EF-1	
1989	Eremothecium gossypii	Database	tuf (EF-1	
1990	Eremothecium gossypii Eremothecium gossypii	Database	tuf (EF-1	
1991	Aspergillus oryzae	Database	tuf (EF-1	
1992	Aureobasidium pullulans	Database	tuf (EF-1	
1993	Histoplasma capsulatum	Database	tuf (EF-1	
1994	Neurospora crassa	Database	tuf (EF-1	
1995	Podospora anserina	Database	tuf (EF-1	
1996	Podospora curvicolla	Database	tuf (EF-1	
1997	Sordaria macrospora	Database	tuf (EF-1	
1998	Trichoderma reesei	Database	tuf (EF-1	
2004	Candida albicans	Database	tuf (M)	
2005	Schizosaccharomyces pombe	Database	tuf (M)	
2010	Klebsiella pneumoniae	Database	blaTEM	
2010	Klebsiella pneumoniae	Database	blaTEM	
2013	Kluyvera ascorbata	This patent	gyrA	
2013	Kluyvera georgiana	This patent	gyrA	
2017	Streptococcus pneumoniae	Database	pbp1A	
2048	Streptococcus pneumoniae	Databas	pbp1A	
2049	Streptococcus pneumoniae	Database	pbp1A	

Table 7. Origin of the nucleic acids and/or sequence is ting (continued).

SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	G ne*
2050	Streptococcus pneumoniae	Database	pbp1A
2051	Streptococcus pneumoniae	Database	pbp1A
2052	Streptococcus pneumoniae	Database	pbp1A
2053	Streptococcus pneumoniae	Database	pbp1A
2054			
2055	Streptococcus pneumoniae	Database	gyrA
	Streptococcus pneumoniae	Database	parC
2056	Streptococcus pneumoniae	This patent	pbp1A
2057	Streptococcus pneumoniae	This patent	pbp1A
2058	Streptococcus pneumoniae	This patent	pbp1A
2059	Streptococcus pneumoniae	This patent	pbp1A
2060	Streptococcus pneumoniae	This patent	pbp1A
2061	Streptococcus pneumoniae	This patent	pbp1A
2062	Streptococcus pneumoniae	This patent	pbp1A
2063	Streptococcus pneumoniae	This patent	pbp1A
2064	· Streptococcus pneumoniae	This patent	pbp1A
2072	Mycobacterium tuberculosis	Database	гроВ
2097	Mycoplasma pneumoniae	Database	tuf
2101	Mycobacterium tuberculosis	Database	inhA
2105	Mycobacterium tuberculosis	Database	embB
2129	Clostridium difficile	Database	cdtA
2130	Clostridium difficile	Database	cdtB
2137	Pseudomonas putida	Genome project	tuf
2138	Pseudomonas aeruginosa	Genome project	tuf
2139	Campylobacter jejuni	Database	atpD
2140	Streptococcus pneumoniae	Database	pbp1a
2144	Staphylococcus aureus	Database	mupA
2147	Escherichia coli	Database	catl
2150	Escherichia coli	Database	catli
2153	Shigella flexneri	Database	catlll
2156	Clostridium perfringens	Database	catP
2159	Staphylococcus aureus	Database	cat
2162	Staphylococcus aureus	Database	cat
2165	Salmonella typhimurium	Database	ppflo-like
2183	Alcaligenes faecalis subsp. faecalis	This patent	tuf
2184	Campylobacter coli	This patent	fusA
2185	Succinivibrio dextrinosolvens	This patent	tuf
2186	Tetragenococcus halophilus	This patent	tuf
2187	Campylobacter jejuni subsp. jejuni	This patent	fusA
2188			fusA
	Campylobacter jejuni subsp. jejuni	This patent	
2189	Leishmania guyanensis	This patent	atpD
2190	Trypanosoma brucei brucei	This patent	atpD
2191	Aspergillus nidulans	This patent	atpD
2192	Leishmania panamensis	This patent	atpD
2193	Aspergillus nidulans	This patent	tuf (M)
2194	Aureobasidium pullulans	This patent	tuf (M)
2195	Emmonsia parva	This patent	tuf (M)
2196	Exserohilum rostratum	This patent	tuf (M)
2197	Fusarium moniliforme	This patent	tuf (M)
2198	Fusarium solani	This patent	tuf (M)
2199	Histoplasma capsulatum	This patent	tuf (M)
2200	Kocuria kristinae	This patent	tuf
2201	Vibrio mimicus	This patent	tuf
2202	Citrobacter freundii	This patent	recA
2203	Clostridium botulinum	This patent	recA
2204	Francisella tularensis	This patent	recA
2205	Peptostreptococcus anaerobius	This patent	recA
2206	Peptostreptococcus asaccharolyticus	This patent	recA
2207	Providencia stuartii	This patent	recA

Table 7. Origin of th nucleic acids and/or sequenc s in the s quenc listing (continued).

SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	Gene*
2208	Salmonella choleraesuis subsp. choleraesuis		
	serotype Paratyphi A	This patent	recA
2209	Salmonella choleraesuis subsp. choleraesuis	The parent	
	serotype Typhimurium	This patent	recA
2210	Staphylococcus saprophyticus	This patent	recA
2211	Yersinia pseudotuberculosis	This patent	recA
2212	Zoogloea ramigera	This patent	recA
2214	Abiotrophia adiacens	This patent	fusA
2215	Acinetobacter baumannii	This patent	fusA
2216	Actinomyces meyeri	This patent	fusA
2217	Clostridium difficile	This patent	fusA
2218	Corynebacterium diphtheriae	This patent	fusA
2219	Enterobacter cloacae	This patent	fusA
2220	Klebsiella pneumoniae subsp. pneumoniae	This patent	fusA
2221	•	This patent	fusA
2222	Listeria monocytogenes	•	fusA
	Mycobacterium avium	This patent	_
2223	Mycobacterium gordonae	This patent	fusA
2224	Mycobacterium kansasii	This patent	fusA
2225	Mycobacterium terrae	This patent	fusA
2226	Neisseria polysaccharea	This patent	fusA
2227	Staphylococcus epidermidis	This patent	fusA
2228	Staphylococcus haemolyticus	This patent	fusA
2229	Succinivibrio dextrinosolvens	This patent	fusA
2230	Tetragenococcus halophilus	This patent	fusA
2231	Veillonella parvula	This patent	fusA
2232	Yersinia pseudotuberculosis	This patent	fusA
2233	Zoogloea ramigera	This patent	fusA
2234	Aeromonas hydrophila	This patent	fusA
2235	Abiotrophia adiacens	This patent	fusA-tuf spac
2236	Acinetobacter baumannii	This patent	fusA-tuf spac
2237	Actinomyces meyeri	This patent	fusA-tuf spac
2238	Clostridium difficile	This patent	fusA-tuf spac
2239	Corynebacterium diphtheriae	This patent	fusA-tuf spac
2240	Enterobacter cloacae	This patent	fusA-tuf spac
2241	Klebsiella pneumoniae subsp. pneumoniae	This patent	fusA-tuf spac
2242	Listeria monocytogenes	This patent	fusA-tuf spac
2243	Mycobacterium avium	This patent	fusA-tuf spac
2244	Mycobacterium gordonae	This patent	fusA-tuf space
2245	Mycobacterium kansasii	This patent	fusA-tuf space
2246	Mycobacterium terrae	This patent	fusA-tuf spac
2247	Neisseria polysaccharea	This patent	fusA-tuf space
2248	Staphylococcus epidermidis	This patent	fusA-tuf space
2249	Staphylococcus haemolyticus	This patent	fusA-tuf spac
2255	Abiotrophia adiacens	This patent	tuf
2256	Acinetobacter baumannii	This patent	tuf
2257	Actinomyces meyeri	This patent	tuf
2258		This patent	tuf
	Clostridium difficile	This patent	tuf
2259	Corynebacterium diphtheriae		tuf
2260	Enterobacter cloacae	This patent	
2261	Klebsiella pneumoniae subsp. pneumoniae	This patent	tuf
2262	Listeria monocytogenes	This patent	tuf
2263	Mycobacterium avium	This patent	tuf
2264	Mycobacterium gordonae	This patent	tuf
2265	Mycobacterium kansasii	This patent	tuf
2266	Mycobacterium terrae	This patent	tuf
2267	Neisseria polysaccharea	This pat nt	tuf
2268	Staphylococcus epidermidis	This patent	tuf
2269	Staphylococcus haemolyticus	This patent	tuf
2270	Aeromonas hydrophila	This pat nt	tuf
2271	Bilophila wadsworthia	This patent	tuf
2272	Brevundimonas diminuta	This patent	tuf
2273	Streptococcus mitis	This patent	pbp1a

Table 7. Origin of the nucleic acids and/or s quences in the sequence listing (continue d).

SEQ ID NO.	Archaeal, bacterial, fungal or parasitical species	Source	G ne*
2274	Streptococcus mitis	This patent	pbp1a
2275	Streptococcus mitis	This patent	pbp1a
2276	Streptococcus oralis	This patent	pbp1a
2277	Escherichia coli	This patent	gyrA
2278	Escherichia coli	This patent	gyrA
2279	Escherichia coli	This patent	gyrA
2280	Escherichia coli	This patent	gyrA
2288	Enterococcus faecium	Database	ddl
2293	Enterococcus faecium	Database	vanA
2296	Enterococcus faecalis	Database	vanB

^{*} tuf indicates tuf sequences, tuf (C) indicates tuf sequences divergent from main (usually A and B) copies of the elongation factor-Tu, tuf (EF-1) indicates tuf sequences of the eukaryotic type (elongation factor 1a), tuf (M) indicates tuf sequences from organellar (mostly mitochondrial) origin.

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fusA indicates fusA sequences; fusA-tuf spacer indicates the intergenic region between fusA and tuf.

atpD indicates atpD sequences of the F-type, atpD (V) indicates atpD sequences of the V-type.

recA indicates recA sequences, recA(Rad51) indicates rad51 sequences or homologs and recA(Dmc1) indicates dmc1 sequences or homologs.

Table 8. Bacterial sp cies us d to test the specificity of the Streptococcus agalactiae-sp clfic amplification primers derived from tuf sequences.

Strain I	Reference number	Strain P	eference numb
Streptococcus acidominimus	ATCC 51726	Bacteroides caccae	ATCC 43185
Streptococcus agalactiae	ATCC 12403	Bacteroides vulgatus	ATCC 8482
Streptococcus agalactiae	ATCC 12973	Bacteroides fragilis	ATCC 25285
Streptococcus agalactiae	ATCC 13813	Candida albicans	ATCC 11006
Streptococcus agalactiae	ATCC 27591	Clostridium innoculum	ATCC 14501
Streptococcus agalactiae	CDCs 1073	Clostridium ramosum	ATCC 25582
Streptococcus anginosus	ATCC 27335	Lactobacillus casei subsp. case	i ATCC 393
Streptococcus anginosus	ATCC 33397	Clostridium septicum	ATCC 12464
Streptococcus bovis	ATCC 33317	Corynebacterium cervicis	NCTC 10604
Streptococcus anginosus	ATCC 27823	Corynebacterium genitalium	ATCC 3303
Streptococcus cricetus	ATCC 19642	Corynebacterium urealyticum	ATCC 4304
Streptococcus cristatus	ATCC 51100	Enterococcus faecalis	ATCC 2921:
Streptococcus downei	ATCC 33748	Enterococcus faecium	ATCC 1943
Streptococcus dysgalactiae	ATCC 43078	Eubacterium lentum	ATCC 4305
Streptococcus equi subsp. equi	ATCC 9528	Eubacterium nodutum	ATCC 3309
Streptococcus ferus	ATCC 33477	Gardnerella vaginalis	ATCC 1401
Streptococcus gordonii	ATCC 10558	Lactobacillus acidophilus	ATCC 4356
Streptococcus macacae	ATCC 35911	Lactobacillus crispatus	ATCC 3382
Streptococcus mitis	ATCC 49456	Lactobacillus gasseri	ATCC 3332
Streptococcus mutans	ATCC 25175	Lactobacillus johnsonii	ATCC 3320
Streptococcus oralis	ATCC 35037	Lactococcus lactis subsp. lactis	
Streptococcus parasanguinis	ATCC 15912	Lactococcus lactis subsp. lactis	ATCC 1145
Streptococcus parauberis	DSM 6631	Listeria innocua	ATCC 3309
Streptococcus pneumoniae	ATCC 27336	Micrococcus luteus	ATCC 934
Streptococcus pyogenes	ATCC 19615	Escherichia coli	ATCC 2592
Streptococcus ratti	ATCC 19645	Micrococcus lylae	ATCC 2756
Streptococcus salivarius	ATCC 7073	Porphyromonas asaccharolytic	a ATCC 2526
Streptococcus sanguinis	ATCC 10556	Prevotella corporis	ATCC 3354
Streptococcus sobrinus	ATCC 27352	Prevotella melanogenica	ATCC 2584
Streptococcus suis	ATCC 43765	Staphylococcus aureus	ATCC 1330
Streptococcus uberis	ATCC 19436	Staphylococcus epidermidis	ATCC 1499
Streptococcus vestubularis	ATCC 49124	Staphylococcus saprophyticus	ATCC 1530

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Table 9. Bact rial speci s used to test the specificity of the *Streptococcus agalactiae*-sp cific amplification primers derived fr m atpD sequ nces.

Strain	Reference number	Strain	Reference number
Streptococcus acidominimus	ATCC 51726	Streptococcus gordonii	ATCC 10558
Streptococcus agalactiae	ATCC 12400	Streptococcus macacae	ATCC 35911
Streptococcus agalactiae	ATCC 12403	Streptococcus mitis	ATCC 49456
Streptococcus agalactiae	ATCC 12973	Streptococcus mutans	ATCC 25175
Streptococcus agalactiae	ATCC 13813	Streptococcus oralis	ATCC 35037
Streptococcus agalactiae	ATCC 27591	Streptococcus parasanguinis	ATCC 15912
Streptococcus agalactiae	CDCs-1073	Streptococcus parauberis	DSM 6631
Streptococcus anginosus	ATCC 27335	Streptococcus pneumoniae	ATCC 27336
Streptococcus anginosus	ATCC 27823	Streptococcus pyogenes	ATCC 19615
Streptococcus bovis	ATCC 33317	Streptococcus ratti	ATCC 19645
Streptococcus cricetus	ATCC 19642	Streptococcus salivarius	ATCC 7073
Streptococcus cristatus	ATCC 51100	Streptococcus sanguinis	ATCC 10556
Streptococcus downei	ATCC 33748	Streptococcus sobrinus	ATCC 27352
Streptococcus downer Streptococcus dysgalactiae	ATCC 43078	Streptococcus suis	ATCC 43765
Streptococcus equi subsp. equ		Streptococcus uberis	ATCC 19436
Streptococcus ferus	ATCC 33477	Streptococcus vestibularis	ATCC 49124

Table 10. Bact rial sp cies used to test the specificity of the *Enterococcus*-sp cific amplification primers dorived from tuf sequence s.

St	rain	Reference number	Strain F	leference numbe
Gram-positiv	e species (n=74))		
Abiotrophia a	diacens	ATCC 49176	Listeria innocua	ATCC 3309
Abiotrophia de		ATCC 49175	Listeria ivanovii	ATCC 1911
Bacillus cereu		ATCC 14579	Listeria mónocytogenes	ATCC 1531
Bacillus subtil	is	ATCC 27370	Listeria seeligeri	ATCC 3596
	m adolescentis	ATCC 27534	Micrococcus luteus	ATCC 9341
Bifidobacteriu	m breve	ATCC 15700	Pediococcus acidilacti	ATCC 3331
Bifidobacteriu	m dentium	ATCC 27534	Pediococcus pentosaceus	ATCC 3331
Bifidobacteriu	m longum	ATCC 15707	Peptococcus niger	ATCC 2773
Clostridium p	•	ATCC 3124	Peptostreptococcus anaerob	ius ATCC 2733
Clostridium se		ATCC 12464	Peptostreptococcus indolicus	s ATCC 2924
	ium aquaticus	ATCC 14665	Peptostreptococcus micros	ATCC 3327
Corynebacter		ATCC 10700	Propionibacterium acnes	ATCC 691
pseudodiphth			Staphylococcus aureus	ATCC 4330
Enterococcus		ATCC 14025	Staphylococcus capitis	ATCC 2784
	casseliflavus	ATCC 25788	Staphylococcus epidermidis	ATCC 1499
Enterococcus		ATCC 43199	Staphylococcus haemolyticu	s ATCC 299
Enterococcus		ATCC 51263	Staphylococcus hominis	ATCC 2784
Enterococcus		ATCC 51266	Staphylococcus lugdunensis	ATCC 438
Enterococcus		ATCC 19432	Staphylococcus saprophytic	
Enterococcus		ATCC 29212	Staphylococcus simulans	ATCC 278
Enterococcus		ATCC 19434	Staphylococcus warneri	ATCC 278
Enterococcus		ATCC 49996	Streptococcus agalactiae	ATCC 138
Enterococcus		ATCC 49573	Streptococcus anginosus	ATCC 333
Enterococcus		ATCC 8044	Streptococcus bovis	ATCC 333
	malodoratus	ATCC 43197	Streptococcus constellatus	ATCC 278
Enterococcus		ATCC 43186	Streptococcus cristatus	ATCC 511
	s pseudoavium	ATCC 49372	Streptococcus intermedius	ATCC 273
Enterococcus		ATCC 49427	Streptococcus mitis	ATCC 494
	saccharolyticus	ATCC 43076	Streptococcus mitis	ATCC 363
Enterococcus		ATCC 49428	Streptococcus mutans	ATCC 271
Enterococcus		ATCC 49903	Streptococcus parasanguini	
Eubacterium		ATCC 49903	Streptococcus pneumoniae	ATCC 277
Gemella hae		ATCC 10379	Streptococcus pneumoniae	ATCC 630
Gemella mor		ATCC 27842	Streptococcus pyogenes	ATCC 196
Lactobacillus		ATCC 4356	Streptococcus salivarius	ATCC 70
	mesenteroides	ATCC 4330 ATCC 19225	Streptococcus sanguinis	ATCC 105
Listeria grayi		ATCC 19223 ATCC 19120	Streptococcus suis	ATCC 437
Listeria grayi Listeria grayi		ATCC 19123	Chiopiooodo odio	7.1.00 407

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Table 10. Bact rial speci s us d to t st the specificity of th Enterococcus-sp cific amplificati n prim rs d rived fr m tuf sequenc s (c ntinued).

Strain	Reference number	Strain I	Reference number		
Gram-negative species (n=39)					
Acidominococcus fermentans	ATCC 2508	Hafnia alvei	ATCC 13337		
Acinetobacter baumannii	ATCC 19606	Klebsiella oxytoca	ATCC 1318		
Alcaligenes faecalis	ATCC 8750	Meganomonas hypermegas			
Anaerobiospirillum	ATCC 29305	Mitsukoella multiacidus	ATCC 2772		
succiniproducens		Moraxella catarrhalis	ATCC 4362		
Anaerorhabdus furcosus	ATCC 25662	Morganella morganii	ATCC 2583		
Bacteroides distasonis	ATCC 8503	Neisseria meningitidis	ATCC 1307		
Bacteroides thetaiotaomicron	ATCC 29741	Pasteurella aerogenes	ATCC 2788		
Bacteroides vulgatus	ATCC 8482	Proteus vulgaris	ATCC 1331		
Bordetella pertussis	LSPQ 3702	Providencia alcalifaciens	ATCC 988		
Bulkholderia cepacia	LSPQ 2217	Providencia rettgeri	ATCC 925		
Butyvibrio fibrinosolvens	ATCC 19171	Pseudomonas aeruginosa	ATCC 2785		
Cardiobacterium hominis	ATCC 15826	Salmonella typhimurium	ATCC 1402		
Citrobacter freundii	ATCC 8090	Serratia marcescens	ATCC 1388		
Desulfovibrio vulgaris	ATCC 29579	Shigella flexneri	ATCC 1202		
Edwardsiellae tarda	ATCC 15947	Shigella sonnei	ATCC 299		
Enterobacter cloacae	ATCC 13047	Succinivibrio dextrinosolvei			
Escherichia coli	ATCC 25922	Tissierella praeacuta	ATCC 255		
Fusobacterium russii	ATCC 25533	Veillonella parvula	ATCC 1079		
Haemophilus influenzae	ATCC 9007	Yersinia enterocolitica	ATCC 961		

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Microbial species for which tuf and/ r atpD and/ r recA s qu nces are available in public databases. Table 11.

	Species	Strain	Accession number	Coding gene
•		tuf sequences		
	Bacteria			
	Actinobacillus actinomycetemcomitans	HK1651	Genome project ²	tuf
	Actinobacillus actinomycetemcomitans	HK1651	Genome project ²	tuf (EF-G)
	Agrobacterium tumefaciens		X99673	tuf
	Agrobacterium tumefaciens		X99673	tuf (EF-G)
	Agrobacterium tumefaciens		X99674	tuf
	Anacystis nidulans	PCC 6301	X17442	tuf tuf
	Aquifex aeolicus	VF5	AE000669	tuf (EF-G)
	Aquifex aeolicus	VF5	AE000669 Genome project ²	tuf (EF-G)
	Aquifex pyrophilus		Y15787	tuf (El -G)
	Aquifex pyrophilus	Ames	Genome project ²	tuf
	Bacillus anthracis	Ames	Genome project ²	tuf (EF-G)
	Bacillus anthracis Bacillus halodurans	C-125	AB017508	tuf
	Bacillus halodurans	C-125	AB017508	tuf (EF-G)
	Bacillus stearothermophilus	CCM 2184	AJ000260	tuf
	Bacillus subtilis	168	D64127	tuf
	Bacillus subtilis	168	D64127	tuf (EF-G)
	Bacillus subtilis	DSM 10	Z99104	tuf
	Bacillus subtilis	DSM 10	Z99104	tuf (EF-G)
	Bacteroides forsythus	ATCC 43037	AB035466	tuf
	Bacteroides fragilis	DSM 1151	- 2	tuf
	Bordetella bronchiseptica	RB50	Genome project ²	tuf
	Bordetella pertussis	Tohama 1	Genome project ²	tuf tuf (EF-G)
	Bordetella pertussis	Tohama 1	Genome project ² U78193	tuf
	Borrelia burdorgferi	B31	AE001155	tuf (EF-G)
	Borrelia burgdorferi	DSM 20425	X76863	tuf
	Brevibacterium linens	Ap	Y12307	tuf
	Buchnera aphidicola	K96243	Genome project ²	tuf (EF-G)
	Burkholderia pseudomallei Campylobacter jejuni	NCTC 11168	Y17167	tuf
	Campylobacter jejuni Campylobacter jejuni	NCTC 11168	CJ11168X2	tuf (EF-G)
	Chlamydia pneumoniae	CWL029	AE001592	tuf
	Chlamydia pneumoniae	CWL029	AE001639	tuf (EF-G)
	Chlamydia trachomatis		M74221	tuf
	Chlamydia trachomatis	D/UW-3/CX	AE001317	tuf (EF-G)
	Chlamydia trachomatis	D/UW-3/CX	AE001305	tuf
	Chlamydia trachomatis	F/IC-Cal-13	L22216	tuf
	Chlorobium vibrioforme	DSM 263	X77033	tuf tuf
	Chloroflexus aurantiacus	DSM 636	X76865	tuf
	Clostridium acetobutylicum	ATCC 824	Genome project ² Genome project ²	tuf
	Clostridium difficile	630 630	Genome project ²	tuf (EF-G)
	Clostridium difficile	NCTC 13129	Genome project ²	tuf
	Corynebacterium diphtheriae	NCTC 13129	Genome project ²	tuf (EF-G)
	Corynebacterium diphtheriae	ASO 19	X77034	tuf
	Corynebacterium glutamicum Corynebacterium glutamicum	MJ-233	E09634	tuf
	Coxiella burnetii	Nine Mile phase		tuf
	Cytophaga lytica	DSM 2039	X77035	tuf
	Deinococcus radiodurans	R1	AE001891	tuf (EF-G)
	Deinococcus radiodurans	R1	AE180092	tuf

Table 11. Microbial species f r which tuf and/ r atpD and/or recA sequences are available in public databases (continued).

Species	Strain	Accession number	Coding gene
Deinococcus radiodurans	R1	AE002041	tuf
	111	1	tuf
Deinonema sp.	ATCC 23834	Z12610	tuf
Eikenella corrodens	ATCC 23834	Z12610	tuf (EF-G)
Eikenella corrodens	A100 23034	Genome project ²	tuf (EF-G)
Enterococcus faecalis		J01690	tuf
Escherichia coli		J01717	tuf
Escherichia coli		X00415	tuf (EF-G)
Escherichia coli		X57091	tuf
Escherichia coli	K-12 MG1655	U00006	tuf
Escherichia coli	K-12 MG1655	U00096	tuf
Escherichia coli		AE000410	tuf (EF-G)
Escherichia coli	K-12 MG1655 DSM 5733	Y15788	tuf
Fervidobacterium islandicum		X76866	tuf
Fibrobacter succinogenes	S85	X76867	tuf
Flavobacterium ferrigeneum	DSM 13524	X59461	tuf
Flexistipes sinusarabici	500 7404	X59461 U09433	tuf
Gloeobacter violaceus	PCC 7421		tuf
Gloeothece sp.	PCC 6501	U09434	tuf
Haemophilus actinomycetemcor	mitans HK1651	Genome project ²	tuf (EF-G)
Haemophilus ducreyi	35000	AF087414	tuf (EF-G)
Haemophilus influenzae	Rd	U32739	tuf
Haemophilus influenzae	Rd	U32746	tuf (EF-G)
Haemophilus influenzae	Rd	U32739	tuf (EP-G)
Helicobacter pylori	26695	AE000511	
Helicobacter pylori	J99	AE001539	tuf (EF-G) tuf
Helicobacter pylori	J99	AE001541	
Herpetosiphon aurantiacus	Hpga1	X76868	tuf Auf
Klebsiella pneumoniae	M6H 78578	Genome project ²	tuf
Klebsiella pneumoniae	M6H 78578	Genome project ²	tuf (EF-G)
Lactobacillus paracasei		E13922	tuf Af
Legionella pneumophila	Philadelphia-1	Genome project ²	tuf
Leptospira interrogans		AF115283	tuf
Leptospira interrogans		AF115283	tuf (EF-G)
Micrococcus luteus	IFO 3333	M17788	tuf (EF-G)
Micrococcus luteus	IFO 3333	M17788	tuf
Moraxella sp.	TAC II 25	AJ249258	tuf
Mycobacterium avium	104	Genome project ²	tuf
Mycobacterium avium	104	Genome project ²	tuf (EF-G)
Mycobacterium bovis	AF2122/97	Genome project ²	tuf
Mycobacterium bovis	AF2122/97	Genome project ²	tuf (EF-G)
Mycobacterium leprae		L13276	tuf
Mycobacterium leprae		Z14314	tuf
Mycobacterium leprae		Z14314	tuf (EF-G)
Mycobacterium leprae	Thai 53	D13869	tuf
Mycobacterium tuberculosis	Erdmann	S40925	tuf
Mycobacterium tuberculosis	H37Rv	AL021943	tuf (EF-G)
Mycobacterium tuberculosis	H37Rv	Z 84395	tuf
Mycobacterium tuberculosis	y42	AD00005	tuf
Mycobacterium tuberculosis	ĆSU#93	Genome project ²	. tuf
Mycobacterium tuberculosis	CSU#93	Genome project ²	tuf (EF-G)
Mycoplasma capricolum	PG-31	X16462	tuf
Mycoplasma genitalium	G37	U39732	tuf
Mycoplasma genitalium	G37	U39689	tuf (EF-G)
Mycoplasma hominis		X57136	tuf
Mycoplasma hominis	PG21	M57675	tuf

Table 11. Micr bial sp cies for which tuf and/or atpD and/ r recA s quences ar availabl in public databases (continu d).

Species	Strain	Accession number	Coding gene
Mycoplasma pneumoniae	M129	AE000019	tuf
Mycoplasma pneumoniae	M129	AE000058	tuf (EF-G)
Neisseria gonorrhoeae	MS11	L36380	tuf
Neisseria gonorrhoeae	MS11	L36380	tuf (EF-G)
Neisseria meningitidis	Z2491	Genome project ²	tuf (EF-G)
Neisseria meningitidis	Z2491	Genome project ²	tuf
Pasteurella multocida	Pm70	Genome project ²	tuf
Peptococcus niger	DSM 20745	X76869	tuf
Phormidium ectocarpi	PCC 7375	U09443	tuf
Planobispora rosea	ATCC 53773	U67308	tuf
Planobispora rosea	ATCC 53733	X98830	tuf
Planobispora rosea	ATCC 53733	X98830	tuf (EF-G)
Plectonema boryanum	PCC 73110	U09444	tuf
Porphyromonas gingivalis	W83	Genome project ²	tuf
Porphyromonas gingivalis	W83	Genome project ²	tuf (EF-G)
Porphyromonas gingivalis	FDC 381	AB035461	tuf
Porphyromonas gingivalis	W83	AB035462	tuf
Porphyromonas gingivalis	SUNY 1021	AB035463	tuf
Porphyromonas gingivalis	A7A1-28	AB035464	tuf
Porphyromonas gingivalis	ATCC 33277	AB035465	tuf
Porphyromonas gingivalis	ATCC 33277	AB035471	tuf (EF-G)
Prochlorothrix hollandica		U09445	tuf
Pseudomonas aeruginosa	PAO-1	Genome project ²	tuf
Pseudomonas putida	• • • •	Genome project ²	tuf
Rickettsia prowazekii	Madrid E	AJ235272	tuf
Rickettsia prowazekii	Madrid E	AJ235270	tuf (EF-G)
Rickettsia prowazekii	Madrid E	Z54171	tuf (EF-G)
Salmonella choleraesuis subsp.			, ,
choleraesuis serotype Typhimurium		X64591	tuf (EF-G)
Salmonella choleraesuis subsp.			,
choleraesuis serotype Typhimurium	LT2 trpE91	X55116	tuf
Salmonella choleraesuis subsp.	,		
choleraesuis serotype Typhimurium	LT2 trpE91	X55117	tuf
Serpulina hyodysenteriae	B204	U51635	tuf
Serratia marcescens		AF058451	tuf
Shewanella putrefaciens	DSM 50426	_1	tuf
Shewanella putrefaciens	MR-1	Genome project ²	tuf
Spirochaeta aurantia	DSM 1902	X76874	tuf
Staphylococcus aureus		AJ237696	tuf (EF-G)
Staphylococcus aureus	EMRSA-16	Genome project ²	tuf
Staphylococcus aureus	NCTC 8325	Genome project ²	tuf
Staphylococcus aureus	COL	Genome project ²	tuf
Staphylococcus aureus	EMRSA-16	Genome project ²	tuf (EF-G)
Stigmatella aurantiaca	DW4	X82820	tuf
Stigmatella aurantiaca	Sg a1	X76870	tuf
Streptococcus mutans	GS-5 Kuramitsu	U75481	tuf
Streptococcus mutans	UAB159	Genome project ²	tuf
Streptococcus oralis	NTCC 11427	P331701	tuf
Streptococcus pyogenes		Genome project ²	tuf (EF-G)
Streptococcus pyogenes	M1-GAS	Genome project ²	tuf
Streptomyces aureofaciens	ATCC 10762	AF007125	tuf
Streptomyces cinnamoneus	Tue89	X98831	tuf
Streptomyces coelicolor	A3(2)	AL031013	tuf (EF-G)
Streptomyces coelicolor	A3(2)	X77039	tuf (EF-G)
Streptomyces coelicolor	M145	X77039	tuf

Table 11. Micr bial sp ci s f r which tuf and/or atpD and/ r recA s quences are available in public databas s (continued).

Species	Strain	Accession number	Coding gene*
Streptomyces collinus	BSM 40733	S79408	tuf
Streptomyces netropsis	Tu1063	AF153618	tuf
Streptomyces ramocissimus		X67057	tuf
Streptomyces ramocissimus		X67058	tuf
Streptomyces ramocissimus		X67057	tuf (EF-G)
Synechococcus sp.	PCC 6301	X17442	tuf (EF-G)
Synechococcus sp.	PCC 6301	X17442	tuf
Synechocystis sp.	PCC 6803	D90913	tuf (EF-G)
Synechocystis sp. Synechocystis sp.	PCC 6803	D90913	tuf (E. G.)
Synechocystis sp.	PCC 6803	X65159	tuf (EF-G)
Taxeobacter occealus	Myx 2105	X77036	tuf (E. C)
	IVIYX 2 103	Genome project ²	tuf (EF-G)
Thermotoga maritima		M27479	tuf (El -G)
Thermotoga maritima	EP 00276	X66322	tuf
Thermus aquaticus		X16278	tuf (EF-G)
Thermus thermophilus	HB8	X05977	tuf
Thermus thermophilus	HB8		
Thermus thermophilus	HB8	X06657	tuf
Thiomonas cuprina	DSM 5495	U78300	tuf
Thiomonas cuprina	DSM 5495	U78300	tuf (EF-G)
Thiomonas cuprina	Hoe5	X76871	tuf
Treponema denticola		Genome project ²	tuf
Treponema denticola		Genome project ²	tuf (EF-G)
Treponema pallidum		AE001202	tuf
Treponema pallidum		AE001222	tuf (EF-G)
Treponema pallidum		AE001248	tuf (EF-G)
Ureaplasma urealyticum	ATCC 33697	Z34275	tuf
Ureaplasma urealyticum	serovar 3 biovar 1		tuf
Ureaplasma urealyticum	serovar 3 biovar 1		tuf (EF-G)
Vibrio cholerae	N16961	Genome project ²	tuf
Wolinella succinogenes	DSM 1740	X76872	tuf
Yersinia pestis	CO-92	Genome project ²	tuf
Yersinia pestis	CO-92	Genome project ²	tuf (EF-G)
Archaebacteria			
Archaeoglobus fulgidus		Genome project ²	tuf (EF-G)
Halobacterium marismortui		X16677	tuf
Methanobacterium thermoautrophicum	delta H	AE000877	tuf
Methanococcus jannaschii	ATCC 43067	U67486	tuf
Methanococcus vannielii		X05698	tuf
Pyrococcus abyssi	Orsay	AJ248285	tuf
Thermoplasma acidophilum	DSM 1728	X53866	tuf
Fungi			
Absidia glauca	CBS 101.48	X54730	<i>tuf</i> (EF-1)
Arxula adeninivorans	Ls3	Z47379	tuf (EF-1)
	KBN616	AB007770	tuf (EF-1)
Aspergillus oryzae	R106	U19723	tuf (EF-1)
Aureobasidium pullulans		Genom project ²	tuf (EF-1)
Candida albicans	SC5314 SC5314	M29934	tuf (EF-1)
Candida albicans	SC5314 SC5314	M29935	tuf (EF-1)
Candida albicans Cryptococcus neoformans	B3501	U81803	tuf (EF-1)
CAVIDATETE DECIONANS	DOOD I	U0 10U3	144 (1.1.7.1)

Table 11. Micr bial speci s for which tuf and/or atpD and/or recA s qu nces ar availabl in public databas s (c ntinu d).

Species	Strain	Accession number	Coding gene
Cryptococcus neoformans	M1-106	U81804	tuf (EF-1)
Eremothecium gossypii	ATCC 10895	X73978	tuf (EF-1)
	A100 10000	A29820	tuf (EF-1)
Eremothecium gossypii	NRRL 26037	AF008498	tuf (EF-1)
Fusarium oxysporum	186AS	U14100	tuf (EF-1)
Histoplasma capsulatum	10043	X74799	tuf (EF-1)
Podospora anserina	M M	X96614	tuf (EF-1)
Podospora curvicolla	VLV	AJ245645	tuf (EF-1)
Prototheca wickerhamii	263-11		tuf (EF-1)
Puccinia graminis	race 32	X73529	tuf (M)
Reclinomonas americana	ATCC 50394	AF007261	
Rhizomucor racemosus	ATCC 1216B	X17475	tuf (EF-1)
Rhizomucor racemosus	ATCC 1216B	J02605	tuf (EF-1)
Rhizomucor racemosus	ATCC 1216B	X17476	tuf (EF-1)
Rhodotorula mucilaginosa		AF016239	tuf (EF-1)
Saccharomyces cerevisiae		K00428	tuf (M)
Saccharomyces cerevisiae		M59369	tuf (EF-G)
Saccharomyces cerevisiae		X00779	tuf (EF-1)
Saccharomyces cerevisiae		X01638	tuf (EF-1)
Saccharomyces cerevisiae		M10992	tuf (EF-1)
Saccharomyces cerevisiae	Alpha S288	X78993	tuf (EF-1)
Saccharomyces cerevisiae		M15666	tuf (EF-1)
Saccharomyces cerevisiae		Z35987	tuf (EF-1)
	S288C (AB972)	U51033	tuf (EF-1)
Saccharomyces cerevisiae	1-40	X94913	tuf (EF-1)
Schizophyllum commune	972h-	AL021816	tuf (EF-1)
Schizosaccharomyces pombe	972h-	AL021813	tuf (EF-1)
Schizosaccharomyces pombe	972h-	D82571	tuf (EF-1)
Schizosaccharomyces pombe	9/211-	U42189	tuf (EF-1)
Schizosaccharomyces pombe	DD745	D89112	tuf (EF-1)
Schizosaccharomyces pombe	PR745		tuf (EF-1)
Sordaria macrospora	000	X96615	tuf (EF-1)
Trichoderma reesei	QM9414	Z23012	tuf (EF-1)
'Yarrowia lipolytica		AF054510	w (EF-1)
Parasites			
Blastocystis hominis	HE87-1	D64080	tuf (EF-1)
Cryptosporidium parvum	. 0.40	U69697	tuf (EF-1)
Eimeria tenella	LS18	A1755521	tuf (EF-1) tuf (EF-1)
Entamoeba histolytica	HM1:IMSS	X83565	
Entamoeba histolytica	NIH 200	M92073	tuf (EF-1)
Giardia lamblia		D14342	tuf (EF-1)
Kentrophoros sp.		AF056101	tuf (EF-1)
Leishmania amazonensis	IFLA/BR/67/PH8		<i>tuf</i> (EF-1)
Leishmania braziliensis		U72244	<i>tuf</i> (EF-1)
Onchocerca volvulus		M64333	tuf (EF-1)
Porphyra purpurea	Avonport	U08844	tuf (EF-1)
Plasmodium berghei	ANKA	AJ224150	tuf (EF-1)
Plasmodium falciparum	K1	X60488	tuf (EF-1)
Plasmodium knowlesi	line H	AJ224153	tuf (EF-1)
	RH	Y11431	tuf (EF-1)
Toxoplasma gondii	ATCC 30207	D78479	tuf (EF-1)
Trichomonas tenax	LVH/75/	U10562	tuf (EF-1)
Trypanosoma brucei		O 10302	10/ (4. 1)
	USAMRU-K/18	L76077	tuf (EF-1)
Trypanosoma cruzi	Υ		

Table 11. Microbial species f r which tuf and/ r atpD and/ r recA sequ nces ar availabl in public databas s (continued).

	Species	Strain	Accession number	Coding gene*
	Human and plants			
	Arabidopsis thaliana	Columbia	X89227	tuf (EF-1)
	Glycine max	Ceresia	X89058	tuf (EF-1)
	Glycine max	Ceresia	Y15107	tuf (EF-1)
	Glycine max	Ceresia	Y15108	tuf (EF-1)
	Glycine max	Maple Arrow	X66062	tuf (EF-1)
	Homo sapiens	•	X03558	<i>tuf</i> (EF-1)
	Pyramimonas disomata		AB008010	tuf
		atpD seque	ences	
	Bacteria			
	Acetobacterium woodi	DSM 1030	U10505	atpD
	Actinobacillus actinomycetemcomitans	HK1651	Genome project ²	atpD
	Bacillus anthracis	Ames	Genome project ²	atpD
	Bacillus firmus	OF4	M60117	atpD
	Bacillus megaterium	QM B1551	M20255	atpD
	Bacillus stearothermophilus		D38058	atpD
	Bacillus stearothermophilus	IFO1035	D38060	atpD
	Bacillus subtilis	168	Z28592	atpD
	Bacteroides fragilis	DSM 2151	M22247	atpD
	Bordetella bronchiseptica	RB50	Genome project ²	atpD
	Bordetella pertussis	Tohama 1	Genome project ²	atpD
	Borrelia burgdorferi	B31	AE001122	atpD (V)
	Burkholderia cepacia	DSM50181	X76877	atpD
)	Burkholderia pseudomallei	K96243	Genome project ²	atpD atpD
	Campylobacter jejuni	NCTC 11168	CJ11168X1	atpD (V)
	Chlamydia pneumoniae	MaD-	Genome project ²	atpD (V)
	Chlamydia trachomatis	MoPn DeM 262	Genome project ² X76873	atpD (V)
_	Chlorobium vibrioforme	DSM 263	AF037156	atpD atpD
5	Citrobacter freundii	JEO503	Genome project ²	atpD atpD
	Clostridium acetobutylicum	ATCC 824	AF101055	atpD
	Clostridium acetobutylicum	DSM 792	Genome project ²	atpD atpD
	Clostridium difficile	630 NCTC13129	Genome project ²	atpD
	Corynebacterium diphtheriae	ASO 19	X76875	atpD
)	Corynebacterium glutamicum	MJ-233	E09634	atpD
	Corynebacterium glutamicum	เหม-233 DSM 2039	M22535	atpD
	Cytophaga lytica	DSM 30053	_3	atpD
	Enterobacter aerogenes	V583	Genome project ²	atpD (V)
	Enterococcus faecalis	¥ J0J	M90060	atpD
•	Enterococcus hirae	ATCC 9790	D17462	atpD (V)
	Enterococcus hirae Escherichia coli	A100 3130	J01594	atpD
	Escherichia coli Escherichia coli		M25464	atpD
			V00267	atpD
	Escherichia coli		V00311	atpD
)	Escherichia coli	K12 MG1655	L10328	atpD
	Escherichia coli	DSM 13524	.3	atpD
	Flavobacterium ferrugineum Haemophilus actinomycetemcomitans	DOM 10024	Genome project ²	atpD
	Haemophilus influenzae	Rd	U32730	atpD
	nacinophilius innucitzae	NCTC 11638	AF004014	atpD

Table 11. Microbial speci s for which tuf and/ r atpD and/or recA sequences ar available in public databases (c ntinu d).

Species	Strain	Accession number	Coding gene
Helicobacter pylori	26695	Genome project ²	atpD
Helicobacter pylori	J99	Genome project ²	atpD
Klebsiella pneumoniae	M6H 78578	Genome project ²	atpD
Lactobacillus casei	DSM 20021	X64542	atpD
Legionella pneumophila	Philadelphia-1	Genome project ²	atpD
Moorella thermoacetica	ATCC 39073	U64318	atpD
Mycobacterium avium	104	Genome project ²	atpD
Mycobacterium bovis	AF2122/97	Genome project ²	atpD
Mycobacterium leprae	711 2 1 2 2 0 7	U15186	atpD
Mycobacterium leprae		Genome project ²	atpD
Mycobacterium tuberculosis	H37Rv	Z73419	atpD
Mycobacterium tuberculosis	CSU#93	Genome project ²	atpD
Mycoplasma gallisepticum	000#30	X64256	atpD
Mycoplasma genitalium	G37	U39725	atpD
	M129	U43738	atpD
Mycoplasma pneumoniae Neisseria gonorrhoeae	FA 1090	Genome project ²	atpD
Neisseria gonormoeae Neisseria meningitidis	Z2491	Genome project ²	atpD
Pasteurella multocida	Pm70	Genome project ²	atpD
	DSM 20465	X64543	atpD
Pectinatus frisingensis	DSM 20475	X76878	atpD
Peptococcus niger	IFAM 1313	X57204	atpD
Pirellula marina	W83	Genome project ²	atpD (V)
Porphyromonas gingivalis	DSM 2376	· X58461	atpD (*)
Propionigenium modestum	PAO1	Genome project ²	atpD
Pseudomonas aeruginosa	PAUI	Genome project ²	atpD
Pseudomonas putida	B100	X99599	atpD
Rhodobacter capsulatus	Б100	X02499	atpD
Rhodospirillum rubrum	F-12	AF036246	atpD
Rickettsia prowazekii		Genome project ²	atpD
Rickettsia prowazekii	Madrid 7ATCC	AB006151	atpD
Ruminococcus albus	JEO4162	AF037155	atpD
Salmonella bongori	BR1859	AF037153	atpD
Salmonella bongori		AF037134 AF037146	atpD
Salmonella choleraesuis	S83769	AFU37 146	aipU
subsp. arizonae		A E0074 47	ntoD
Salmonella choleraesuis	u24	AF037147	atpD
subsp. arizonae	1/000	AE027140	atnD
Salmonella choleraesuis subsp.	K228	AF037140	atpD
choleraesuis serotype Dublin	1/774	AE007400	atpD
Salmonella choleraesuis subsp.	K771	AF037139	aipu
choleraesuis serotype Dublin	Divoc oc	AE027140	atpD
Salmonella choleraesuis subsp.	Div36-86	AF037142	aipu
choleraesuis serotype Infantis	D: 05 00	AE0074.40	ataD
Salmonella choleraesuis subsp.	Div95-86	AF037143	atpD
choleraesuis serotype Tennessee	1.70	AE007444	otoD)
Salmonella choleraesuis subsp.	LT2	AF037141	atpD
choleraesuis serotype Typhimurium	0004040	AF0074.40	ete.D
Salmonella choleraesuis	DS210/89	AF037149	atpD
subsp. diarizonae		45000110	e+-0
Salmonella choleraesuis	JEO307	AF037148	atpD
subsp. diarizonae		A 500 m 1 50	- 4- D
Salmonella choleraesuis	S109671	AF037150	atpD
subsp. <i>diarizonae</i>		.=	
Salmonella choleraesuis	S84366	AF037151	atpD
subsp. houtenae	42.3		
Salmonella choleraesuis	S84098	AF037152	atpD

Table 11. Microbial species f r which tuf and/or atpD and/or recA sequenc s ar availabl in public databases (continued).

Species	Strain	Accession number	Coding gene
subsp. houtenae			
Salmonella choleraesuis	BR2047	AF037153	atpD
subsp. indica	DITEOTI	711 007 100	4.70
Salmonella choleraesuis	NSC72	AF037144	atpD
subsp. <i>salamae</i>	113072	A1 007 144	aipo
	S114655	AF037145	atpD
Salmonella choleraesuis	3114033	AI 037 149	aipo
subsp. salamae	MD 4	Conomo project ²	atpD
Shewanella putrefaciens	MR-1	Genome project ²	
Staphylococcus aureus	COL	Genome project ²	atpD
Stigmatella aurantiaca	Sga1	X76879	atpD
Streptococcus bovis	JB-1	AB009314	atpD
Streptococcus mutans	GS-5	U31170	atpD
Streptococcus mutans	UAB159	Genome project ²	atpD
Streptococcus pneumoniae	Type 4	Genome project ²	atpD (V)
Streptococcus pneumoniae	Type 4	Genome project ²	atpD
Streptococcus pyogenes	M1-GAS	Genome project ²	atpD (V)
Streptococcus pyogenes	M1-GAS	Genome project ²	atpD
Streptococcus sanguinis	10904	AF001955	atpD
Streptomyces lividans	1326	Z22606	atpD
Thermus thermophilus	HB8	D63799	atpD (V)
Thiobacillus ferrooxidans	ATCC 33020	M81087	atpD
Treponema pallidum	Nichols	AE001228	atpD (V)
Vibrio alginolyticus		X16050	atpD
Vibrio cholerae	N16961	Genome project ²	atpD
Wolinella succinogenes	DSM 1470	X76880	atpD
Yersinia enterocolitica	NCTC 10460	AF037157	atpD
Yersinia pestis	CO-92	Genome project ²	atpD
Archaebacteria			
Archaeoglobus fulgidus	DSM 4304	AE001023	atpD (V)
Halobacterium salinarum		S56356	atpD (V)
Haloferax volcanii	WR 340	X79516	atpD
Methanococcus jannaschii	DSM 2661	U67477	atpD (V)
Methanosarcina barkeri	DSM 800	J04836	atpD (V)
Fungi		-	
	005044	0	c4mD
Candida albicans	SC5314	Genome project ²	atpD
Candida tropicalis	0050450	M64984	atpD (V)
Kluyveromyces lactis	2359/152	U37764	atpD
Neurospora crassa		X53720	atpD
Saccharomyces cerevisiae		M12082	atpD
Saccharomyces cerevisiae	X2180-1A	J05409	atpD (V)
Schizosaccharomyces pombe	972 h-	\$47814	atpD (V)
Schizosaccharomyces pombe	972 h-	M57956	atpD
Parasit s			
			A . 5
Giardia lamblia	WB	U18938	atpD
Plasmodium falciparum	3D7	L08200	atpD (V)
Trypanosoma congolense	IL3000	Z25814	atpD (V)

Table 11. Microbial species f r which tuf and/or atpD and/or recA s qu nces ar availabl in public databas s (continu d).

	public databas s (continu d).					
	Species	Strain	Accession number	Coding gene		
	Human and plants					
	Homo sapiens		L09234	atpD (V)		
	Homo sapiens		M27132	atpD		
		recA seque	nces			
	Bacteria					
	Acetobacter aceti	no. 1023	S60630	recA		
	Acetobacter altoacetigenes	MH-24	E05290	recA		
	Acetobacter polyoxogenes	NBI 1028	D13183	recA		
	Acholeplasma laidlawii	8195	M81465	recA		
	Acidiphilium facilis	ATCC 35904	D16538	recA		
	Acidothermus cellulolyticus	ATCC 43068	AJ006705	recA		
	Acinetobacter calcoaceticus	BD413/ADP1	L26100	recA		
	Actinobacillus actinomycetemcomitans	HK1651	Genome project ²	recA		
	Aeromonas salmonicida	A449	U83688	recA		
	Agrobacterium tumefaciens	C58	L07902	recA		
	Allochromatium vinosum	\ <i>/</i> EE	AJ000677	recA		
	Aquifex aeolicus	VF5	AE000775	recA		
	Aquifex pyrophilus	Kol5a	L23135	recA		
	Azotobacter vinelandii	40	S96898	recA		
	Bacillus stearothermophilus	10	Genome project ²	recA recA		
	Bacillus subtilis	PB1831	U87792			
	Bacillus subtilis	168	Z99112	recA		
)	Bacteroides fragilis	NOED OOFO	M63029	recA recA		
	Bifidobacterium breve	NCFB 2258	AF094756	recA recA		
	Blastochloris viridis	DSM 133	AF022175	recA recA		
	Bordetella pertussis	165	X53457			
	Bordetella pertussis	Tohama I	Genome project ²	recA		
5	Borrelia burgdorferi	Sh-2-82	U23457	recA		
	Borrelia burgdorferi	B31	AE001124	recA		
	Brevibacterium flavum	MJ-233	E10390	recA		
	Brucella abortus	2308	L00679	recA		
	Burkholderia cepacia	ATCC 17616	U70431	recA		
)	Burkholderia cepacia	1/00040	D90120	recA		
	Burkholderia pseudomallei	K96243	Genome project ²	recA		
	Campylobacter fetus subsp. fetus	23D	AF020677	recA		
	Campylobacter jejuni	81-176	U03121	recA		
	Campylobacter jejuni	NCTC 11168	AL139079	recA		
5	Chlamydia trachomatis	12	U16739	recA		
	Chlamydia trachomatis	D/UW-3/CX	AE001335	recA		
	Chlamydophila pneumoniae	CWL029	AE001658	recA		
	Chloroflexus aurantiacus	J-10-fl	AF037259	recA		
	Clostridium acetobutylicum	4.0	M94057	recA		
)	Clostridium perfringens	13	U61497	recA		
	Corynebacterium diphtheriae	NCTC13129	Genome project ²	recA		
	Corynebacterium glutamicum	AS019	U14965	recA		
	Corynebacterium pseudotuberculosis	C231	U30387	recA		
	Deinococcus radiodurans	KD8301	AB005471	recA		
5	Deinococcus radiodurans	R1	U01876	recA		

Table 11. Microbial species for which *tuf* and/or *atpD* and/ r *recA* s qu nces are available in public databases (c ntinued).

Species	Strain	Accession number	Coding gene
Enterobacter agglomerans	339	L03291	recA
Enterococcus faecalis			recA
Erwinia carotovora			recA
Escherichia coli			recA
Escherichia coli			recA
Escherichia coli	K-12	AE000354	recA
Frankia alni	OGIX M81466 X55554 J01672 X55552 K-12 AF000354 Arl3 AJ006707 U21001 Rd U32687 Rd U32741 Rd L07529 69A Z35478 26695 AE000536 J99 AE001453 M6H 78578 Genome project ² ML3 M88106 X55453 serovar patoc Serovar pomona U29169 MS-1 MS-1 MFK1 M35325 ATCC 31226 X59514 104 Genome project ² X73822 H37Rv X58485 CSU#93 Genome project ² X78822 H37Rv X58485 CSU#93 Genome project ² X78822 H37Rv X58485 CSU#93 Genome project ² X78822 H37Rv X58485 CSU#93 Genome project ² X73822 H37Rv X58485 CSU#93 Genome project ² X73822 H37Rv X58485 CSU#93 Genome project ² X73826 H37Rv X58485 CSU#93 Genome project ² X73822 H37Rv X58485 CSU#93 Genome project ² X73822 H37Rv X58485 CSU#93 Genome project ² X73822 H37Rv X58485 CSU#93 Genome project ² AF122/97 ATCC 29342 MPAE000033 KD735 L22074 L40368 L40367 NCTC 10212 L57910 LCDC 81-176 AJ223869 LNP 1646 U57906 NCTC 10294 AJ223871 Vedros M601 AJ223870 CCUG 2131 AJ223882 CCUG 4165A AJ223881 NCTC 11050 AJ223878 NHITCC 2376 AJ223879 Bangor 9 AJ223873 LNP 4444 U57907 CH95 FA19 X64842	recA	
Gluconobacter oxydans		U21001	recA
Haemophilus influenzae	Rd	U32687	recA
Haemophilus influenzae	Rd	U32741	recA
Haemophilus influenzae	Rd	L07529	recA
Helicobacter pylori	69A	Z35478	recA
Helicobacter pylori	26695	AE000536	recA
Helicobacter pylori	J99	AE001453	recA
Klebsiella pneumoniae	M6H 78578	Genome project ²	recA
Lactococcus lactis	ML3		recA
Legionella pneumophila		X55453	recA
Leptospira biflexa	serovar patoc	U32625	recA
Leptospira interrogans	· ·	U29169	recA
Magnetospirillum magnetotacticum		X17371	recA
Methylobacillus flagellatus	MFK1	M35325	recA
Methylomonas clara	ATCC 31226	X59514	recA
Mycobacterium avium	104	Genome project ²	recA
Mycobacterium bovis	AF122/97	Genome project ²	recA
Mycobacterium leprae		X73822	recA
Mycobacterium tuberculosis	H37Rv		recA
Mycobacterium tuberculosis	CSU#93	Genome project ²	recA
Mycoplasma genitalium	G37	U39717	recA
Mycoplasma mycoides	GM9	L22073	recA
Mycoplasma pneumoniae	ATCC 29342	MPAE000033	recA
Mycoplasma pulmonis	KD735	L22074	recA
Myxococcus xanthus		L40368	recA
Myxococcus xanthus		L40367	recA
Neisseria animalis	NCTC 10212	U57910	recA
Neisseria cinerea	LCDC 81-176	AJ223869	recA
Neisseria cinerea	LNP 1646		recA
Neisseria cinerea			recA
Neisseria cinerea			recA
Neisseria elongata			recA ·
Neisseria elongata	NHITCC 2376		recA
Neisseria elongata	CCUG 4557	AJ223879	recA
subsp. intermedia			
Neisseria flava			recA
Neisseria flavescens			recA
Neisseria gonorrhoeae			recA
Neisseria gonorrhoeae			recA
Neisseria gonorrhoeae	MS11	X17374	recA
Neisseria gonorrhoeae		Genome project ²	recA
Neisseria lactamica	CCUC 7757	AJ223866	recA
Neisseria lactamica	CCUG 7852	Y11819	recA
Neisseria lactamica	LCDC 77-143	Y11818	recA
Neisseria lactamica	LCDC 80-111	AJ223864	recA

Table 11. Micr bial species for which *tuf* and/or *atpD* and/or *recA* sequ nces ar availabl in public databases (continued).

Species	Strain	Accession number	Coding gene
Neisseria lactamica	LCDC 845	AJ223865	recA
Neisseria lactamica	NCTC 10617	U57905	recA
Neisseria lactamica	NCTC 10618	AJ223863	recA
Neisseria meningitidis	44/46	X64849	recA
Neisseria meningitidis	Bangor 13	AJ223868	recA
Neisseria meningitidis	HF116	X64848	recA
Neisseria meningitidis Neisseria meningitidis	HF130	X64844	recA
Neisseria meningitidis	HF46	X64847	recA
		·	
Neisseria meningitidis	M470	X64850	recA
Neisseria meningitidis	N94II	X64846	recA
Neisseria meningitidis	NCTC 8249	AJ223867	recA
Neisseria meningitidis	P63	X64845	recA
Neisseria meningitidis	S3446	U57903	recA
Neisseria meningitidis	FAM18	Genome project ²	recA
Neisseria mucosa	LNP 405	U57908	гесА
Neisseria mucosa	Vedros M1801	AJ223875	recA
Neisseria perflava	CCUG 17915	AJ223876	recA
Neisseria perflava	LCDC 85402	AJ223862	recA
Neisseria pharyngis var. flava	NCTC 4590	U57909	recA
Neisseria polysaccharea	CCUG 18031	Y11815	recA
Neisseria polysaccharea	CCUG 24845	Y11816	recA
Neisseria polysaccharea	CCUG 24846	Y11814	recA
Neisseria polysaccharea	INS MA 3008	Y11817	recA
	NCTC 11858	U57904	recA
Neisseria polysaccharea		AJ223872	recA
Neisseria sicca	NRL 30016		
Neisseria subflava	NRL 30017	AJ223874	recA
Paracoccus denitrificans	DSM 413	U59631	recA
Pasteurella multocida		X99324	recA
Porphyromonas gingivalis	W83	U70054	recA
Prevotella ruminicola	JCM 8958	U61227	recA
Proteus mirabilis	pG1300	X14870	recA
Proteus vulgaris		X55555	recA
Pseudomonas aeruginosa		X05691	recA
Pseudomonas aeruginosa	PAM 7	X52261	recA
Pseudomonas aeruginosa	PAO12	D13090	recA
Pseudomonas fluorescens	OE 28.3	M96558	recA
Pseudomonas putida		L12684	recA
Pseudomonas putida	PpS145	U70864	recA
Rhizobium leguminosarum	VF39	X59956	recA
biovar <i>viciae</i>	V1 05	700000	100/1
	CNPAF512	X62479	recA
Rhizobium phaseoli		X82183	
Rhodobacter capsulatus	J50		recA
Rhodobacter sphaeroides	2.4.1	X72705	recA
Rhodopseudomonas palustris	N 7	D84467	recA
Rickettsia prowazekii	Madrid E	AJ235273	recA
Rickettsia prowazekii	Madrid E	U01959	recA
Serratia marcescens		M22935	recA
Shigella flexneri		X55553	recA
Shigella sonnei	KNIH104S	AF101227	recA
Sinorhizobium meliloti	2011	X59957	recA
Staphylococcus aureus		L25893	recA
Streptococcus gordonii	Challis V288	L20574	гесА
Streptococcus mutans	UA96	M81468	recA
Streptococcus mutans Streptococcus mutans	GS-5	M61897	recA
on epicococas maians	~ 0-5.	11101007	,6071

Table 11. Microbial sp cles f r which tuf and/or atpD and/or recA s quences are available in public databases (c ntinu d).

Species	Strain	Acc ssion number	Coding gene*
Streptococcus pneumoniae	R800	Z34303	recA
Streptococcus pyogenes	NZ131	U21934	recA
Streptococcus pyogenes	D471	M81469	recA
Streptococcus salivarius	•	M94062	recA
subsp. thermophilus			
Streptomyces ambofaciens	DSM 40697	Z30324	recA
Streptomyces coelicolor	A3(2)	AL020958	recA .
Streptomyces lividans	TK24	X76076	recA
Streptomyces rimosus	R6	X94233	recA
Streptomyces venezuelae	ATCC10712	U04837	recA
Synechococcus sp.	PR6	M29495	recA
Synechocystis sp.	PCC6803	D90917	recA
Thermotoga maritima		L23425	recA
Thermotoga maritima		AE001823	recA
Thermus aquaticus		L20095	recA
Thermus thermophilus	HB8	D17392	recA
Thiobacillus ferrooxidans		M26933	recA
Treponema denticola		Genome project ²	recA
Treponema pallidum	Nichols	AE001243	recA
Vibrio anguillarum		M80525	recA
Vibrio cholerae	017	X71969	recA
Vibrio cholerae	2740-80	U10162	recA
Vibrio cholerae	569B	L42384	recA
Vibrio cholerae	M549	AF117881	recA
Vibrio cholerae	M553	AF117882	recA
Vibrio cholerae	M645	AF117883	recA
Vibrio cholerae	M793	AF117878	recA
Vibrio cholerae	M794	AF117880	recA
Vibrio cholerae	M967	AF117879	recA
Xanthomonas citri	XW47	AF006590	recA
Xanthomonas oryzae		AF013600	recA
Xenorhabdus bovienii	T228/1	U87924	recA
Xenorhabdus nematophilus	AN6	AF127333	recA
Yersinia pestis	231	X75336	recA
Yersinia pestis	CO-92	Genome project ²	recA
Fungi, parasites, human and plan	ts		
Anabaena variabilis	ATCC 29413	M29680	recA
Arabidopsis thaliana		U43652	recA (Rad5
Candida albicans		U39808	recA (Dmc1
Coprinus cinereus	Okayama-7	U21905	recA (Rad5
Emericella nidulans		Z80341	recA (Rad5
Gallus gallus		L09655	recA (Rad5
Homo sapiens		D13804	recA (Rad5
Homo sapiens		D63882	recA (Dmc1
Leishmania major	Friedlin	AF062379	recA (Rad5
Leishmania major	Friedlin	AF062380	recA (Dmc1
Mus musculus		D58419	recA (Dmc1
Neurospora crassa	74-OR23-1A	D29638	recA (Rad5
Saccharomyces cerevisiae		D10023	recA (Rad5
Schizosaccharomyces pombe	•	Z22691	recA (Rad5
Schizosaccharomyces pombe	972h-	AL021817	recA (Dmc1
Tetrahymena thermophila	PB9R	AF064516	recA (Rad5

Tabl 11. Microbial species f r which tuf and/ r atpD and/or recA sequenc s are available in public databases (continued).

Species	Strain	Accession number	Coding gene*
Trypanosoma brucei	stock 427	Y13144	recA (Rad51)
Ustilago maydis		U62484	recA (Rad51)
Xenopus laevis		D38488	recA (Rad51)
Xenopus laevis		D38489	recA (Rad51)

^{*} tuf indicates tuf sequences, including tuf genes, fusA genes and fusA-tuf intergenic spacers.
tuf (C) indicates tuf sequences divergent from main (usually A and B) copies of the elongation factor-Tu
tuf (EF-1) indicates tuf sequences of the eukaryotic type (elongation factor 1α)
tuf (M) indicates tuf sequences from organellar (mostly mitochondrial) origin
atpD indicates atpD sequences of the F-type
atpD (V) indicates atpD sequences of the V-Type
recA indicates recA sequences
recA (Rad51) indicates rad51 sequences or homologs

5

10

15

recA (Dmc1) indicates dmc1 sequences or homologs

¹ Nucleotides sequences published in Arch. Microbiol. 1990 **153**:241-247
² These sequences are from theTIGR database (http://www.tigr.org/tdb/tdb.html)

³ Nucleotides sequences published in FEMS Microbiology Letters 1988 **50**:101-106

Table 12. Bacterial species used to t st th specificity of the *Staphylococcus*-specific amplification prim rs deriv d fr m *tuf* s quenc s.

Stra	ain	Reference number	Strain R	eference number
Stanhylococc	al species (n=27)	Other Gram-positive bacter	ia (n=20)
			Bacillus subtilis	ATCC 27370
Staphylococcu Staphylococcu	s aureus	ATCC 43957 ATCC 35844	Enterococcus avium	ATCC 14025
subsp. anaero. Staphylococcu		ATCC 43300	Enterococcus durans	ATCC 19432
subsp. aureus	S aureus	A100 40000	Emeroodod darans	
Staphylococcu	s auricularis	ATCC 33753	Enterococcus faecalis	ATCC 19433
Staphylococcu		ATCC 27840	Enterococcus faecium	ATCC 19434
subsp. capitis	•			
Staphylococcu		ATCC 35538	Enterococcus flavescens	ATCC 49996
Staphylococcu		ATCC 51365	Enterococcus gallinarum	ATCC 49573
	is chromogenes	ATCC 43764	Lactobacillus acidophilus	ATCC 4356
Staphylococcu		DSM 20260	Lactococcus lactis	ATCC 11454
subsp. urealyti		ATOO 40474	t into sin immonso	ATCC 3309
Staphylococcu		ATCC 49171	Listeria innocua	ATCC 33090
Staphylococcu		ATCC 14990	Listeria ivanovii	ATCC 1511
Staphylococcu		ATCC 43958	Listeria monocytogenes Macrococcus caseolyticus	ATCC 1354
Staphylococcu		ATCC 49168 ATCC 35539	Streptococcus agalactiae	ATCC 1381
Staphylococcu		ATCC 35539 ATCC 29970	Streptococcus agaiactiae Streptococcus anginosus	ATCC 3339
	ıs haemolyticus	ATCC 29970 ATCC 27844	Streptococcus anginosus Streptococcus bovis	ATCC 3331
Staphylococcu Staphylococcu		ATCC 27844 ATCC 11249	Streptococcus mutans	ATCC 2517
	ıs intermedius	ATCC 29663	Streptococcus pneumoniae	ATCC 630
Staphylococcu		ATCC 43959	Streptococcus pyogenes	ATCC 1961
Staphylococcu		ATCC 29070	Streptococcus salivarius	ATCC 707
	ıs lugdunensis	ATCC 43809	O Op. 1000000000000000000000000000000000000	
	us saprophyticus	ATCC 15305		
Staphylococci		ATCC 49545		
subsp. coagul				
Staphylococci		ATCC 29060		
subsp. sciuri				
Staphylococci	us simulans	ATCC 27848		
Staphylococci	us warneri	ATCC 27836		
Staphylococci	us xylosus	ATCC 29971		
Gram-negativ	/e bacteria (n=33	3)		
Acinetobacter	baumannii	ATCC 19606	Morganella morganii	ATCC 2583
Bacteroides d		ATCC 8503	Neisseria gonorrhoeae	ATCC 3520
Bacteroides fr	ragilis	ATCC 25285	Neisseria meningitidis	ATCC 1307
Bulkholderia d		ATCC 25416	Proteus mirabilis	ATCC 2593
Bordetella per	rtussis	ATCC 9797	Proteus vulgaris	ATCC 133
Citrobacter fre	eundii	ATCC 8090	Providencia rettgeri	ATCC 925
Enterobacter	aerogenes	ATCC 13048	Providencia stuartii	ATCC 299
Enterobacter		ATCC 13047	Pseudomonas aeruginosa	ATCC 2785
Escherichia c		ATCC 25922	Pseudomonas fluorencens	ATCC 1352
Haemophilus		ATCC 8907	Salmonella choleraesuis	ATCC 1400
	parahaemolyticus		Salmonella typhimurium	ATCC 810
•	parainfluenzae	ATCC 7901	Serratia marcescens	ATCC 810 ATCC 120
Hafnia alvei		ATCC 13337	Shigella flexneri	ATCC 120
Kingella indol	-	ATCC 25869	Shigella sonnei	
Klebsiella oxy		ATCC 13182	Stenotrophomonas maltoph Yersinia enterocolitica	ATCC 136
Klebsiella pne	eumoniae	ATCC 13883	reisinia enterocontica	A100 301

Table 13. Bact rial sp cies us d to test the sp cificity of the penicillin-r sistant *Streptococcus pneumoniae* assay.

	Strain	Reference number	Strain Re	eference numb
G	ram-positive species (n=67)			
	biotrophia adiacens	ATCC 49175	Staphylococcus hominis	ATCC 2784
Al	biotrophia defectiva	ATCC 49176	Staphylococcus lugdunensis	ATCC 4380
A	ctinomyces pyogenes	ATCC 19411	Staphylococcus saprophyticus	ATCC 1530
Ba	acillus anthracis	ATCC 4229	Staphylococcus simulans	ATCC 2784
Ba	acillus cereus	ATCC 14579	Staphylococcus. warneri	ATCC 2783
Bi	ifidobacterium breve	ATCC 15700	Streptococcus acidominimus	ATCC 5172
C	lostridium difficile	ATCC 9689	Streptococcus agalactiae	ATCC 1240
E	nterococcus avium	ATCC 14025	Streptococcus anginosus	ATCC 3339
E	nterococcus casseliflavus	ATCC 25788	Streptococcus bovis	ATCC 3331
E	nterococcus dispar	ATCC 51266	Streptococcus constellatus	ATCC 2782
	nterococcus durans	ATCC 19432	Streptococcus cricetus	ATCC 1962
E	nterococcus faecalis	ATCC 29212	Streptococcus cristatus	ATCC 5110
E	nterococcus faecium	ATCC 19434	Streptococcus downei	ATCC 3374
E	nterococcus flavescens	ATCC 49996	Streptococcus dysgalactiae	ATCC 4307
E	nterococcus gallinarum	ATCC 49573	Streptococcus equi	ATCC 952
	nterococcus hirae	ATCC 8043	Streptococcus ferus	ATCC 3347
E	nterococcus mundtii	ATCC 43186	Streptococcus gordonii	ATCC 1059
E	nterococcus raffinosus	ATCC 49427	Streptococcus intermedius	ATCC 273
Lá	actobacillus lactis	ATCC 19435	Streptococcus mitis	ATCC 90
Lá	actobacillus monocytogenes	ATCC 15313	Streptococcus mitis	LSPQ 258
	lobiluncus curtisii	ATCC 35242	Streptococcus mitis	ATCC 494
P	eptococcus niger	ATCC 27731	Streptococcus mutans	ATCC 2717
	eptostreptococcus acones	ATCC 6919	Streptococcus oralis	ATCC 105
	eptostreptococcus anaerobius	ATCC 27337	Streptococcus oralis	ATCC 981
	eptostreptococcus	ATCC 2639	Streptococcus oralis	ATCC 350
	saccharolyticus		Streptococcus parasanguinis	ATCC 159
	eptostreptococcus lactolyticus	ATCC 51172	Streptococcus parauberis	ATCC 663
	eptostreptococcus magnus	ATCC 15794	Streptococcus rattus	ATCC 159
	eptostreptococcus prevotii	ATCC 9321	Streptococcus salivarius	ATCC 707
	eptostreptococcus tetradius	ATCC 35098	Streptococcus sanguinis	ATCC105
	taphylococcus aureus	ATCC 25923	Streptococcus suis	ATCC 437
	taphylococcus capitis	ATCC 27840	Streptococcus uberis	ATCC 194
	taphylococcus epidermidis	ATCC 14990	Streptococcus vestibularis	ATCC 491
	taphylococcus haemolyticus	ATCC 29970		
	ram-negative species (n=33)			
A	ctinetobacter baumannii	ATCC 19606	Moraxella morganii	ATCC 130
B	ordetella pertussis	ATCC 9797	Neisseria gonorrhoeae	ATCC 352
C	itrobacter diversus	ATCC 27028	Neisseria meningitidis	ATCC 130
C	itrobacter freundii	ATCC 8090	Proteus mirabilis	ATCC 259
E	nterobacter aerogenes	ATCC 13048	Proteus vulgaris	ATCC 133
	nterobacter agglomerans	ATCC 27155	Providencia alcalifaciens	ATCC 988
	nterobacter cloacae	ATCC 13047	Providencia rettgeri	ATCC 925
E	scherichia coli	ATCC 25922	Providencia rustigianii	ATCC 336
Н	laemophilus ducreyi	ATCC 33940	Providencia stuartii	ATCC 336
	laemophilus haemolyticus	ATCC 33390	Pseudomonas aeruginosa	ATCC 355
	laemophilus influenzae	ATCC 9007	Pseudomonas fluorescens	ATCC 135
	laemophilus parainfluenzae	ATCC 7901	Pseudomonas stutzeri	ATCC 175
	lafnia alvei	ATCC 13337	Salmonella typhimurium	ATCC 140
	lebsiella oxytoca	ATCC 13182	Serratia marcescens	ATCC 138
	lebsiella pneumoniae	ATCC 13883	Shigella flexneri	ATCC 120
K	iebsielia priedifibrilae	71.00 10000		
	foraxella atlantae	ATCC 29525	Yersina enterocolitica	ATCC 961

Table 14. Bacterial sp cies (n=104) det ct d by th plat let contaminants assay. Bold characters indicate the major bacterial contaminants found in platelet concentrates.

•					Stanbulanceus simulans
5	Abiotrophia adiacens		Klebsiella oxytoca		Staphylococcus simulans
	Abiotrophia defectiva		Klebsiella pneumoniae		Staphylococcus warneri Stenotrophomonas maltophilia
	Acinetobacter baumannii		Legionella pneumophila	00	
	Acinetobacter Iwoffi		Megamonas hypermegale	80	Streptococcus acidominimus
	Aerococcus viridans	45	Moraxella atlantae		Streptococcus agalactiae
0	Bacillus anthracis		Moraxella catarrhalis		Streptococcus anginosus
	Bacillus cereus		Morganella morganii		Streptococcus bovis
	Bacillus subtilis		Neisseria gonorrheae		Streptococcus constellatus
	Brucella abortus		Neisseria meningitidis	85	Streptococcus cricetus
	Burkholderia cepacia	50	Pasteurella aerogenes		Streptococcus cristatus
5	Citrobacter diversus		Pasteurella multocida		Streptococcus dysgalactiae
	Citrobacter freundii		Peptostreptococcus magnus		Streptococcus equi
	Enterobacter aerogenes		Proteus mirabilis	•	Streptococcus ferus
	Enterobacter agglomerans		Providencia alcalifaciens	90	Streptococcus gordonii
	Enterobacter cloacae	55	Providencia rettgeri		Streptococcus intermedius
0	Enterococcus avium		Providencia rustigianii		Streptococcus macacae
•	Enterococcus casseliflavus		Providencia stuartii		Streptococcus mitis
	Enterococcus dispar		Pseudomonas aeruginosa		Streptococcus mutans
	Enterococcus durans		Pseudomonas fluorescens	95	Streptococcus oralis
	Enterococcus faecalis	60	Pseudomonas stutzeri		Streptococcus parasanguinis
25	Enterococcus faecium		Salmonella bongori		Streptococcus parauberis
-	Enterococcus flavescens		Salmonella choleraesuis		Streptococcus pneumoniae
	Enterococcus gallinarum		Salmonella enteritidis		Streptococcus pyogenes
	Enterococcus mundtii		Salmonella gallinarum	100	Streptococcus ratti
	Enterococcus raffinosus	65			Streptococcus salivarius
0	Enterococcus solitarius		Serratia liquefaciens		Streptococcus sanguinis
U	Escherichia coli		Serratia marcescens		Streptococcus sobrinus
	Gemella morbillorum		Shigella flexneri		Streptococcus uberis
	Haemophilus ducreyi		Shigella sonnei	105	Streptococcus vestibularis
	Haemophilus haemolyticus	70			Vibrio cholerae
. 5	Haemophilus influenzae	,0	Staphylococcus capitis		Yersinia enterocolitica
35			Staphylococcus epidermidis		Yersinia pestis
	Haemophilus		Staphylococcus haemolyticus		Yersinia pseudotuberculosi
	parahaemolyticus		Staphylococcus hominis		
	Haemophilus parainfluenzae	7.5	Stanhylococcus Hominis		

75 Staphylococcus lugdunensis

Staphylococcus saprophyticus

Hafnia alvei

40 Kingella kingae

Table 15. Microorganism entified by commercial systems¹.

Acinetobacter introductors (phins) Acinetobacter introductors (phins) Acinetobacter individual (Acinetobacter introductors) Acinetobacter addiresistens Acinetobacter addiresistens Acinetobacter addiresistens Acinetobacter addiresistens Acinetobacter species Acinetobacter addiresistens Acinetobactius perceivante Acinetobactius perceivante Acinetobacter addiresistens Acinetobactius perceivante Acinetoproses naesundii 35 Acinetoryces naesundii 35 Acinetoryces naesundii 36 Acinetoryces naesundii Acinetoryces naesundii 37 Acinetoryces naesundii Acinetoryces naesundii Acinetoryces naesundii 38 Acinetoryces naesundii Acinetoryces properes Acinetoryces propereivante Acinetoryces propereiv						
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Actionabacter apociles Acidaminococcus termantans Acinatobacter alcalegenes Acinatobacter calcaceaticus biovar Acinatobacter calcaceaticus biovar Acinatobacter promocilis Acinatobacter portionia Acinatobacter promocilis Acinatobactilis seguiui Acinatobactilis acinatomycalamocomilis Acinatobactilis acinatomycalamocomilis Acinatobactilis acinatomycalamocomilis Acinatobactilis acinatomycalamocomilis Acinatobactilis acinatomycalamocomilis Acinatobactilis seguiui Acinatobactilis acinatomycalimocomilis Acinatobactilis seguiui Acinatobactilis seguiu		adjacens)				
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Actinomyces species Actinomyces unicensis Actinomyces turicensis Actinomyces turicensis Actinomyces viscosus Aerococcus species Aerococcus species Aerococcus viridans Aeromonas hydrophila Aeromonas hydrophila group Aeromonas jandaei Aeromonas salmonicida Aeromonas salmonicida subsp. achmongenes Aeromonas selmonicida subsp. achmongenes Aeromonas selmonicida subsp. achmonida aubsp. Bacteroides species Bacteroides species Bacteroides splanchnicus Bact						
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Actinomyces viscosus Bacteroides vatus Bacteroides splanchnicus Bacteroides splanchnicus Bacteroides splanchnicus Bacteroides splanchnicus Candida (Clavispora) lusitaniae Candida (Pichia) guilliemondii Candida (Cardida Didnii) Candida catenulata Candida cellicules Candida dellicules Candida dellicules Candida dellicules Candida dellicules Candida dellicules Candida prose Candida dellicules Candida dellicules Candida dellicules Candida dellicules Candida dellicules Candida prose Candida pros	40					
Aerococcus species Aerococcus species Aerococcus species Aeromonas caviae Aeromonas hydrophila Aeromonas hydrophila group Aeromonas salmonicida Aeromonas salmonicida subsp. Bacteroides uritomis Bacte			115		190	Campylobacter upsaliensis
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Aeromonas schubertii Aeromonas sobria Aeromonas species Aeromonas trota Bifidobacterium infantis Aeromonas trota Bifidobacterium infantis Aeromonas veronii Aeromonas veronii biovar sobria Aeromonas veronii biovar veronii Agrobacterium radiobacter Agrobacterium tumefaciens Alcaligenes denitrificans Alcaligenes odorans Alcaligenes odorans Alcaligenes species Alcaligenes species Alcaligenes xylosoxidans Alcaligenes xylos	55					·
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Aeromonas veronii Aeromonas veronii biovar sobria Aeromonas veronii biovar sobria Aeromonas veronii biovar veronii Agrobacterium radiobacter Agrobacterium species Alcaligenes denitrificans Alcaligenes odorans Alcaligenes odorans Alcaligenes odorans Alcaligenes species Alcaligenes sylosoxidans Alcaligenes xylosoxidans		Aeromonas species				
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Aeromonas veronii biovar suronii Aeromonas veronii biovar veronii Agrobacterium radiobacter Agrobacterium species Bordetella parapertussis Candida ketyr Candida ketyr Candida ketyr Candida krusei Candida magnoliae Candida magnoliae Candida magnoliae Candida magnoliae Candida magnoliae Candida meribiosica Candida meribranaefaciens Candida meribranaefaciens Candida norvegensis Candida norvegensis Candida norvegica Candida norvegica Candida norvegica Candida parapsilosis Candida parapsilosis Candida parapsilosis	60		125			
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Agrobacterium radiobacter Agrobacterium species Bordetella parapertussis Bordetella pertussis Candida mariobacter Candida marioba						
Agrobacterium species 65 Agrobacterium tumefaciens Alcaligenes denitrificans Alcaligenes faecalis Alcaligenes odorans Alcaligenes odorans Alcaligenes odorans Alcaligenes odorans Alcaligenes odorans (Alcaligenes Alcaligenes species Alcaligenes species Alcaligenes species Alcaligenes species Alcaligenes species Alcaligenes xylosoxidans Brevibacterium epidermidis Candida lambica Candida magnoliae Candida magnoli						
Alcaligenes denitrificans Alcaligenes faecalis Alcaligenes odorans Alcaligenes odorans Alcaligenes odorans Alcaligenes odorans (Alcaligenes Alcaligenes odorans (Alcaligenes Alcaligenes species Alcaligenes species Alcaligenes species Alcaligenes xylosoxidans		Agrobacterium species				
Alcaligenes faecalis Alcaligenes odorans Alcaligenes odorans Alcaligenes odorans (Alcaligenes Alcaligenes odorans (Alcaligenes Alcaligenes odorans (Alcaligenes Alcaligenes species Alcaligenes species Alcaligenes species Alcaligenes xylosoxidans	03	Agrobacterium tumetacieris	140			Candida magnoliae
Alcaligenes odorans Alcaligenes odorans Alcaligenes odorans (Alcaligenes Alcaligenes odorans (Alcaligenes Branhamella species Brevibacillus brevis Alcaligenes species Alcaligenes species Alcaligenes xylosoxidans Alcaligenes xylosoxidans Brevibacterium casei Alcaligenes xylosoxidans subsp. Brevibacterium epidermidis Candida melibiosica Candida membranaefaciens Candida norvegensis Candida norvegica Candida parapsilosis Candida melibiosica Candida parapsilosis		Alcaligeries deminincaris	140		215	
Alcaligenes odorans (Alcaligenes Branhamella species Candida membranaelaciens Brevibacillus brevis Candida norvegensis Candida norvegensis Candida norvegensis Candida norvegensis Candida norvegica Candida norvegica Alcaligenes xylosoxidans Brevibacterium casei 220 Candida parapsilosis Candida parapsilosis Candida paratropicalis Candida paratropicalis Candida paratropicalis Candida paratropicalis	,	Alceligenes adorens		Branhamella (Moraxella) catarrhalis		
70 faecalis) Alcaligenes species Alcaligenes xylosoxidans Alcaligenes xylosoxidans Alcaligenes xylosoxidans Brevibacillus laterosporus Brevibacterium casei Candida norvegensis Candida norvegica Candida parapsilosis Candida parapsilosis Candida parapsilosis Candida paratropicalis Brevibacterium epidermidis Candida paratropicalis		Alcalinanes odorans (Alcalinanes				=
Alcaligenes species 145 Brevibacillus laterosporus Candida norvegica Alcaligenes xylosoxidans Brevibacterium casei 220 Candida parapsilosis Alcaligenes xylosoxidans subsp. Brevibacterium epidermidis Candida paratropicalis	76					
Alcaligenes xylosoxidans Brevibacterium casei 220 Candida paratspilosis Alcaligenes xylosoxidans subsp. Brevibacterium epidermidis Candida paratropicalis Candida paratropicalis	,,		145			
Alcaligenes xylosoxidans subsp. Brevibactenum epidermidis Carolida paratropicans		Alcaligenes xylosoxidans			220	
		Alcaligenes xylosoxidans subsp.		Brevibacterium epidermidis		
deninicans biovidacionali inolo		denitrificans		Brevibacterium linens		Candida pelliculosa

Table 15. Microorganism :ntifled by commercial systems (continued)

	Candida pseudotropicalis	00	Clostridium hastiforme		Corynebacterium urealyticum (group
	Candida pulcherrima	80	Clostridium histolyticum		D2)
	Candida ravautii		Clostridium innocuum	140	Corynebacterium xerosis
	Candida rugosa		Clostridium limosum	160	Cryptococcus albidus
	Candida sake		Clostridium novyi		Cryptococcus ater
	Candida silvicola	0.5	Clostridium novyi A		Cryptococcus cereanus
	Candida species	85	Clostridium paraputrificum		Cryptococcus gastricus
	Candida sphaerica		Clostridium perfringens	165	Cryptococcus humicolus
	Candida stellatoidea		Clostridium putrificum	165	Cryptococcus lactativorus
) (Candida tenuis		Clostridium ramosum		Cryptococcus laurentii
f	Candida tropicalis		Clostridium septicum		Cryptococcus luteolus
í	Candida utilis	90	Clostridium sordellii		Cryptococcus melibiosum
- (Candida valida		Clostridium species		Cryptococcus neoformans
	Candida vini		Clostridium sphenoides	170	Cryptococcus species
_	Candida viswanathii		Clostridium sporogenes		Cryptococcus terreus
_	Candida zeylanoides		Clostridium subterminale		Cryptococcus uniguttulatus
	Capnocytophaga gingivalis	95	Clostridium tertium		Debaryomyces hansenii
	Capnocytophaga ochracea	,,,	Clostridium tetani		Debaryomyces marama
	Capnocytophaga species		Clostridium tyrobutyricum	175	Debaryomyces polymorphus
			Comamonas (Pseudomonas)	1,5	Debaryomyces species
	Capnocytophaga sputigena		acidovorans		Demabacter hominis
	Cardiobacterium hominis	100			Dermacoccus (Micrococcus)
	Camobacterium divergens	100	Comamonas (Pseudomonas)		
	Camobacterium piscicola		testosteroni	180	nishinomiyaensis
	CDC group ED-2		Comamonas species	190	The state of the s
	CDC group EF4 (Pasteurella sp.)		Corynebacterium accolens		Edwardsiella hoshinae
	CDC group EF-4A		Corynebacterium afermentans		Edwardsiella ictaluri
	CDC group EF-4B	105	Corynebacterium amycolatum		Edwardsiella species
	CDC group EQ-Z		Corynebacterium aquaticum		Edwardsiella tarda
	CDC group HB-5		Corynebacterium argentoratense	185	Eikenella corrodens
0	CDC group II K-2		Corynebacterium auris		Empedobacter brevis (Flavobacteriu
	CDC group IV C-2 (Bordetella-like)		Corynebacterium bovis		breve)
	CDC group M5	110	Corynebacterium coyleae		Enterobacter aerogenes
	CDC group M6		Corynebacterium cystitidis		Enterobacter agglomerans
	Cedecea davisae		Corynebacterium diphtheriae	190	Enterobacter amnigenus
_	Cedecea lapagei		Corynebacterium diphtheriae biotype		Enterobacter amnigenus asburiae
	Cedecea napage. Cedecea neteri		belfanti		(CDC enteric group 17)
	Cedecea species	115	Corynebacterium diphtheriae biotype		Enterobacter amnigenus biogroup 1
		113	gravis		Enterobacter amnigenus biogroup 2
	Cellulomonas (Oerskovia) turbata		Corynebacterium diphtheriae biotype	195	Enterobacter asburiae
	Cellulomonas species		intermedius	173	Enterobacter cancerogenus
10	Chlamydia species				Enterobacter cloacae
	Chromobacterium violaceum	120	Corynebacterium diphtheriae biotype		
	Chryseobacterium (Flavobacterium)	120	mitis		Enterobacter gergoviae
	indologenes		Corynebacterium flavescens	200	Enterobacter hormaechei
	Chryseobacterium (Flavobacterium)		Corynebacterium glucuronolyticum	200	Enterobacter intermedius
15	meningosepticum		Corynebacterium glucuronolyticum-		Enterobacter sakazakii
	Chryseobacterium gleum		seminale		Enterobacter species
	Chryseobacterium species	125	Corynebacterium group A		Enterobacter taylorae
	Chryseomonas indologenes		Corynebacterium group A-4		Enterobacter taylorae (CDC enteric
	Citeromyces matritensis		Corynebacterium group A-5	205	group 19)
0	Citrobacter amalonaticus		Corynebacterium group ANF		Enterococcus (Streptococcus)
	Citrobacter braakii		Corynebacterium group B		cecorum
	Citrobacter diversus	130	Corynebacterium group B-3		Enterococcus (Streptococcus) faeca
	Citrobacter farmeri		Corynebacterium group F		(Group D)
	Citrobacter freundii		Corynebacterium group F-1	210	
55	Citrobacter freundii complex		Corvnebacterium group F-2		faecium(Group D)
,,,	Citrobacter koseri		Corynebacterium group G		Enterococcus (Streptococcus)
	Citrobacter koseri Citrobacter sedlakii	135	Corynebacterium group G-1		saccharolyticus
		133	Corynebacterium group G-2		Enterococcus avium (Group D)
	Citrobacter species		Corynebacterium group 1	215	
c 10	Citrobacter werkmanii		Corynebacterium group 1-2	210	(Steptococcus faecium subsp.
50	Citrobacter youngae		Corynebacterium jeikeium (group JK)		casseliflavus)
	Clostridium acetobutylicum	1.40			Enterococcus durans (Streptococcu
	Clostridium barati	140			faecium subsp. durans) (Group D)
	Clostridium beijerinckii		murium)	220	
	Clostridium bifermentans		Corynebacterium macginleyi	220	
55	Clostridium botulinum .		Corynebacterium minutissimum		Enterococcus hirae
	Clostridium botulinum (NP) B&F		Corynebacterium pilosum		Enterococcus malodoratus
	Clostridium botulinum (NP) E	145			Enterococcus mundtii
	Clostridium botulinum (P) A&H		Corynebacterium		Enterococcus raffinosus
	Clostridium botulinum (P) F		pseudodiphtheriticum	225	
70	Clostridium botulinum G1		Corynebacterium pseudotuberculosis		Erwinia amylovora
	Clostridium botulinum G2		Corynebacterium pyogenes		Erwinia carotovora
	Clostridium butyricum	150	Corynebacterium renale		Erwinia carotovora subsp. atrosept
		-50	Corynebacterium renale group		Erwinia carotovora subsp.
	Clostridum cedavers				
	Clostridium cadaveris		Corvnebacterium seminala	230	betavasculorum
	Clostridium chauvoei		Corynebacterium seminale	230	
	Clostridium chauvoei Clostridium clostridiiforme		Corynebacterium species	230	Erwinia carotovora subsp. carotovo
	Clostridium chauvoei	155	Corynebacterium species Corynebacterium striatum (C.	230	betavasculorum Erwinia carotovora subsp. carotovo Erwinia chrysanthemi Erwinia cypripedii

Table 15. Microorganism entified by commercial systems (continu d)

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	Erwinia nigrifluens		VII		Lactobacillus paracasei subsp.
	Erwinia riigililaeris Erwinia quercina	80	Haemophilus parainfluenzae biotype		paracasei
	Erwinia rhapontici	00	VIII		Lactobacillus pentosus
	Erwinia rubrifaciens		Haemophilus paraphrohaemolyticus	160	Lactobacillus plantarum
5	Erwinia salicis		Haemophilus paraphrophilus		Lactobacillus salivarius
•	Erwinia species		Haemophilus segnis		Lactobacillus salivarius var. salicinius
	Erysipelothrix rhusiopathiae	85	Haemophilus somnus		Lactobacillus species
	Erysipelothrix species		Haemophilus species		Lactococcus diacitilactis
	Escherichia blattae		Hafnia alvei	165	•
1 0 ·	Escherichia coli		Hanseniaspora guilliermondil		Lactococcus lactis subsp. cremoris
	Escherichia coli A-D	00	Hanseniaspora uvarum		Lactococcus lactis subsp. diacitilactis Lactococcus lactis subsp. hordniae
	Escherichia coli O157:H7	90	Hanseniaspora valbyensis		Lactococcus lactis subsp. lactis
	Escherichia fergusonii		Hansenula anomala	170	Lactococcus plantarum
	Escherichia hermannii		Hansenula holstii Hansenula polymorpha	170	Lactococcus raffinolactis
15	Escherichia species		Helicobacter (Campylobacter) cinaedi		Leciercia adecarboxylata
	Escherichia vulneris Eubacterium aerofaciens	95			Legionella species
	Eubacterium alactolyticum	,,	fennelliae		Leminorella species
	Eubacterium lentum		Helicobacter (Campylobacter) pylori	175	Leptospira species
20	Eubacterium limosum		Issatchenkia orientalis		Leptotrichia buccalis
	Eubacterium species		Kingella denitrificans		Leuconostoc (Weissella)
	Ewingella americana	100	Kingella indologenes		paramesenteroides
	Filobasidiella neoformans		Kingella kingae	• • • •	Leuconostoc camosum
	Filobasidium floriforme		Kingella species	180	
25	Filobasidium uniguttulatum		Klebsiella omithinolytica		Leuconostoc gelidum Leuconostoc lactis
	Flavimonas oryzihabitans	105	Klebsiella oxytoca		Leuconostoc mesenteroides
	Flavobacterium gleum	105			Leuconostoc mesenteroides subsp.
	Flavobacterium indologenes		Klebsiella pneumoniae subsp. ozaenae	185	
20	Flavobacterium odoratum		Klebsiella pneumoniae subsp.	100	Leuconostoc mesenteroides subsp.
30	Flavobacterium species Francisella novicida		pneumoniae		dextranicum
	Francisella philomiragia	110	Klebsiella pneumoniae subsp.		Leuconostoc mesenteroides subsp.
	Francisella species	110	rhinoscleromatis		mesenteroides
	Francisella tularensis		Klebsiella species	190	Leuconostoc species
35			Klebsiella terrigena		Listeria grayi
	Fusobacterium necrogenes		Kloeckera apiculata		Listeria innocua
	Fusobacterium necrophorum	115	Kloeckera apis		Listeria ivanovii
	Fusobacterium nucleatum		Kloeckera japonica	195	Listeria monocytogenes
	Fusobacterium species		Kloeckera species	193	
40	Fusobacterium varium		Kluyvera ascorbata		Listeria seeligeri Listeria species
	Gaffkya species	120	Kluyvera cryocrescens Kluyvera species		Listeria welshimeri
	Gardnerella vaginalis	120	Kluyveromyces lactis		Megasphaera elsdenii
	Gemella haemolysans Gemella morbillorum		Kluyveromyces marxianus	200	
45			Kluyveromyces thermotolerans		Metschnikowia pulchemima
, -3	Geotrichum candidum		Kocuria (Micrococcus) kristinae		Microbacterium species
-	Geotrichum fermentans	125	Kocuria (Micrococcus) rosea		Micrococcus luteus
	Geotrichum penicillarum		Kocuria(Micrococcus) varians	205	Micrococcus Iylae
	Geotrichum penicillatum		Koserella trabulsii	205	
50	Geotrichum species		Kytococcus (Micrococcus) sedentarius		Mobiluncus curtisii
	Gordona species	100	Lactobacillus (Weissella) viridescens		Mobiluncus mulieris Mobiluncus species
	Haemophilus aegyptius	130	Lactobacillus A		Moellerella wisconsensis
	Haemophilus aphrophilus		Lactobacillus acidophilus Lactobacillus B	210	
	Haemophilus ducreyi		Lactobacillus brevis	210	Moraxella atlantae
55			Lactobacillus buchneri		Moraxella bovis
	Haemophilus haemolyticus Haemophilus influenzae	135			Moraxella lacunata
	Haemophilus influenzae biotype l		Lactobacillus casei subsp. casei		Moraxella nonliquefaciens
	Haemophilus influenzae biotype II		Lactobacillus casei subsp. lactosus	215	
60			Lactobacillus casei subsp. mamnosus		Moraxella phenylpyruvica
	Haemophilus influenzae biotype IV		Lactobacillus catenaformis		Moraxella species
	Haemophilus influenzae biotype V	140			Morganella morganii
	Haemophilus influenzae biotype VI		Lactobacillus collinoides	220	Morganella morganii subsp. morganii Morganella morganii subsp. sibonii
_	Haemophilus influenzae biotype VII		Lactobacillus coprophilus	220	Mycobacterium africanum
65			Lactobacillus crispatus		Mycobacterium asiaticum
	Haemophilus paragallinarum	146	Lactobacillus curvatus Lactobacillus delbrueckii subsp.		Mycobacterium avium
	Haemophilus parahaemolyticus	145	bulgaricus deibrueckii suusp.		Mycobacterium bovis
	Haemophilus parainfluenzae		Lactobacillus delbrueckii subsp.	225	Mycobacterium chelonae
70	Haemophilus parainfluenzae biotype I Haemophilus parainfluenzae biotype II		delbrueckii		Mycobacterium fortuitum
/0	Haemophilus parainfluenzae biotype Haemophilus parainfluenzae biotype		Lactobacillus delbrueckii subsp. lactis		Mycobacterium gordonae
	III	150			Mycobacterium kansasii
	Haemophilus parainfluenzae biotype		Lactobacillus fructivorans		Mycobacterium malmoense
	IV		Lactobacillus helveticus	230	
75			Lactobacillus helveticus subsp. jugurti		Mycobacterium phlei
	Haemophilus parainfluenzae biotype		Lactobacillus jensenii		Mycobacterium scrofulaceum
	VI	15:			Mycobacterium smegmatis
	Haemophilus parainfluenzae biotype		Lactobacillus minutus		Mycobacterium species

Table 15. Microorganisms ntified by comm rolal syst ms (continued)¹.

٨	Aycobacterium tuberculosis		Pichia fermentans	•	Saccharomyces exiguus
	Aycobacterium ulcerans	80	Pichia membranaefaciens		Saccharomyces kluyverii
		-	Pichia norvegensis		Saccharomyces species
	Aycobacterium xenopi		Pichia ohmeri	160	Sakaguchia dacryoides
	Aycoplasma fermentans		Pichia spartinae		(Rhodosporidium dacryoidum)
	Aycoplasma hominis				Salmonella arizonae
	Aycoplasma orale	06	Pichia species		Salmonella choleraesuis
	Aycoplasma pneumoniae	85	Plesiomonas shigelloides		Salmonella enteritidis
	Nycoplasma species		Porphyromonas asaccharolytica	165	
	<i>Myroides</i> species		Porphyromonas endodontalis	105	Salmonella gallinarum
f	Neisseria cinerea		Porphyromonas gingivalis		Salmonella paratyphi A
1	Veisseria elongata subsp. elongata		Porphyromonas levii		Salmonella paratyphi B
1	Veisseria flava	90	Prevotella (Bacteroides) buccae		Salmonella pullorum
	Neisseria flavescens		Prevotella (Bacteroides) buccalis		Salmonella species
1	Neisseria gonorrhoeae		Prevotella (Bacteroides) corporis	170	Salmonella typhi
	Veisseria lactamica		Prevotella (Bacteroides) denticola		Salmonella typhimurium
	Neisseria meningitidis		Prevotella (Bacteroides) loescheii		Salmonella typhisuis
	Veisseria mucosa	95			Salmonella/Arizona
	Neisseria perflava		Prevotella (Bacteroides) disiens		Serratia ficaria
	Neisseria polysaccharea		Prevotella (Bacteroides)oris	175	Serratia fonticola
	Neisseria saprophytes		Prevotella bivia (Bacteroides bivius)		Serratia grimesii
	Neisseria sapropriytes Neisseria sicca		Prevotella intermedia (Bacteroides		Serratia liquefacions
		100	intermedius)		Serratia marcescens
	Neisseria subflava	100	Prevotella melaninogenica		Serratia odorifera
- 4	Neisseria weaveri		(Bacteroides melaninogenicus)	180	Serratia odorifera type 1
	Neisseria weaveri (CDC group M5)		Prevotella ruminicola		Serratia odorifera type 2
	Nocardia species		Propionibacterium acnes		Serratia plymuthica
	Ochrobactrum anthropi	105			Serratia proteamaculans
	Oerskovia species	105	Propionibacterium avidum		Serratia proteamaculans subsp.
	Oerskovia xanthineolytica		Propionibacterium granulosum	185	proteamaculans
	Oligella (Moraxella) urethralis		Propionibacterium propionicum	103	Serratia proteamaculans subsp.
	Oligella species		Propionibacterium species		
	Oligella ureolytica		Proteus mirabilis		quinovora
	Paenibacillus alvei	110			Serratia rubidaea
	Paenibacillus macerans		Proteus species	100	Serratia species
	Paenibacillus polymyxa		Proteus vulgaris	190	•
	Pantoea agglomerans		Prototheca species		Alteromonas) putrefaciens
	Pantoea ananas (Erwinia uredovora)		Prototheca wickerhamii		Shigella boydii
	Pantoea dispersa	115	Prototheca zopfii		Shigella dysenteriae
	Pantoea species		Providencia alcalifaciens		Shigella flexneri
	Pantoea stewartii		Providencia heimbachae	195	Shigella sonnei
	Pasteurella (Haemophilus) avium		Providencia rettgeri		Shigella species
			Providencia rustigianii		Sphingobacterium multivorum
	Pasteurella aerogenes	120			Sphingobacterium species
	Pasteurella gallinarum	120	Providencia stuartii		Sphingobacterium spiritivorum
	Pasteurella haemolytica		Providencia stuartii urea +	200	
	Pasteurella haemolyticus		Pseudomonas (Chryseomonas)		Sphingomonas (Pseudomonas)
	Pasteurella multocida				paucimobilis
	Pasteurella multocida SF	125	luteola		Sporidiobolus salmonicolor
	Pasteurella multocida subsp.	125			Sporobolomyces roseus
	multocida		Pseudomonas aeruginosa	205	
	Pasteurella multocida subsp. septica		Pseudomonas alcaligenes	205	Sporobolomyces salmonicolor
	Pasteurella pneumotropica		Pseudomonas cepacia		Sporobolomyces species
	Pasteurella species		Pseudomonas chlororaphis (P.		Staphylococcus (Peptococcus)
	Pasteurella ureae	130			saccharolyticus
	Pediococcus acidilactici		Pseudomonas fluorescens		Staphylococcus arlettae
	Pediococcus damnosus		Pseudomonas fluorescens group	210	Staphylococcus aureus
	Pediococcus pentosaceus		Pseudomonas mendocina		Staphylococcus aureus (Coagula
	Pediococcus species		Pseudomonas pseudoalcaligenes		negative)
	Peptococcus niger	135			Staphylococcus auricularis
	Peptococcus species		Pseudomonas species		Staphylococcus capitis
	Peptococcus species Peptostreptococcus anaerobius		Pseudomonas stutzeri	215	Staphylococcus capitis subsp. ca
	Peptostreptococcus asaccharolyticus		Pseudomonas testosteroni		Staphylococcus capitis subsp.
)			Pseudomonas vesicularis		ureolyticus
	Peptostreptococcus Indolicus	140			Staphylococcus caprae
	Peptostreptococcus magnus	140	alactolyticus		Staphylococcus camosus
	Peptostreptococcus micros		Psychrobacter (Moraxella)	220	
	Peptostreptococcus parvulus				Staphylococcus chromogenes
•			phenylpyruvicus Babaalla egystilis		Staphylococcus cohnii
	Peptostreptococcus productus	1.40	Rahnella aquatilis		Staphylococcus connii subsp. co.
	Peptostreptococcus species	145			Staphylococcus cohnii subsp. Co.
	Peptostreptococcus tetradius		Burkholderia) pickettii	225	
	Phaecoccomyces exophialiae		Rhodococcus (Corynebacterium) equi	225	
)	Photobacterium damselae		Rhodococcus species		Staphylococcus epidermidis
,	Pichia (Hansenula) anomala		Rhodosporidium toruloides		Staphylococcus equorum
,	Pichia (Hansenula) jadinii	150			Staphylococcus gallinarum
,			Rhodotorula minuta	_	Staphylococcus haemolyticus
,	Pichia (Hansenuta) netersonu			230) Staphylococcus hominis
,	Pichia (Hansenula) petersonii Pichia angusta (Hansenula		rinogolorgia muchaginosa (n. 1001a)		
	Pichia angusta (Hansenula		Rhodotorula mucilaginosa (R. rubra) Rhodotorula species		Staphylococcus hominis subsp.
5	Pichia angusta (Hansenula polymorpha)		Rhodotorula species	200	
	Pichia angusta (Hansenula	155	Rhodotorula species Rickettsia species	200	Staphylococcus hominis subsp.

Table 15. Micro rganisms identified by comm rcial syst ms (continued)1.

		60	Streptococcus Gamma (non)-		Tetragenococcus (Pediococcus)
	Charle dangerer building	00	hemolytic	120	halophilus
	Staphylococcus hyicus Staphylococcus intermedius		Streptococcus gordonii		Torulaspora delbrueckii
	Staphylococcus kloosii		Streptococcus Group B		(Saccharomyces rosei)
_	Staphylococcus lentus		Streptococcus Group C		Torulopsis candida
5	Staphylococcus lentus Staphylococcus lugdunensis	65	Streptococcus Group D		Torulopsis haemulonii
	Staphylococcus luguurierisis Staphylococcus saprophyticus	05	Streptococcus Group E	125	Torulopsis inconspicua
			Streptococcus Group F		Treponema species
	Staphylococcus schleiferi		Streptococcus Group G		Trichosporon asahii
10	Staphylococcus sciuri Staphylococcus simulans		Streptococcus Group L		Trichosporon asteroides
10	Staphylococcus species	70			Trichosporon beigelii
	Staphylococcus warneri		Streptococcus Group U	130	Trichosporon cutaneum
	Staphylococcus xylosus		Streptococcus intermedius		Trichosporon inkin
	Stenotrophomonas (Xanthomonas)		Streptococcus intermedius		Trichosporon mucoides
15	maltophilia		(Streptococcus milleri II)		Trichosporon ovoides
13	Stephanoascus ciferrii	75	Streptococcus intermedius (viridans		Trichosporon pullulans
	Stomatococcus mucilaginosus		Streptococcus)	135	Trichosporon species
	Streptococcus acidominimus		Streptococcus milleri group		Turicella otitidis
	Streptococcus agalactiae		Streptococcus mitis		Ureaplasma species
20	Streptococcus agalactiae (Group B)		Streptococcus mitis (viridans		Ureaplasma urealyticum
	Streptococcus agalactiae hemolytic	80			Veillonella parvula (V. alcalescens)
	Streptococcus agalactiae non-		Streptococcus mitis group	140	
	h molytic		Streptococcus mutans		Vibrio alginolyticus
	Streptococcus alactolyticus		Streptococcus mutans (viridans		Vibrio cholerae
25	Streptococcus anginosus	0.5	Streptococcus)		Vibrio damsela Vibrio fluvialis
	Streptococcus anginosus (Group D,	85		145	
	nonenterococci)		Streptococcus parasanguis Streptococcus pneumoniae	145	Vibrio harveyi
	Streptococcus beta-hemolytic group A		Streptococcus priedmoniae Streptococcus porcinus		Vibrio hollisae
20	Streptococcus beta-hemolytic non-		Streptococcus pyogenes		Vibrio metschnikovii
30	group A or B	90			Vibrio mimicus
	Streptococcus beta-hemolytic non-	70	Streptococcus salivarius	150	Vibrio parahaemolyticus
	group A Streptococcus beta-hemolytic		Streptococcus salivarius (viridans		Vibrio species
	Streptococcus bovis (Group D,		Streptococcus)		Vibrio species SF
35	nonenterococci)		Streptococcus salivarius subsp.		Vibrio vulnificus
22	Streptococcus bovis I	95	salivarius		Weeksella (Bergeylla) virosa
	Streptococcus bovis II		Streptococcus salivarius subsp.	155	
	Streptococcus canis		thermophilus		Weeksella virosa
	Streptococcus constellatus		Streptococcus sanguis		Williopsis (Hansenula) satumus
40			Streptococcus sanguis I (viridans		Xanthomonas campestris
	(Streptococcus milleri I)	100		160	Xanthomonas species Yarrowia (Candida) lipolytica
	Streptococcus constellatus (viridans		Streptococcus sanguis II	100	Yersinia aldovae
	Streptococcus)		Streptococcus sanguis II (viridans		Yersinia anterocolitica
	Streptococcus downei		Streptococcus) Streptococcus sobrinus		Yersinia enterocolitica group
45		105			Yersinia frederiksenii
	dysgalactiae Streptococcus dysgalactiae subsp.	10.	Streptococcus suis I .	165	
	equisimilis		Streptococcus suis II		Yersinia intermedius
	Streptococcus equi (Group C/Group G		Streptococcus uberis		Yersinia kristensenii
50			Streptococcus uberis (viridans		Yersinia pestis
50	Streptococcus equi subsp. equi	110) Streptococcus)		Yersinia pseudotuberculosis
	Streptococcus equi subsp.		Streptococcus vestibularis	170	
	zooepidemicus		Streptococcus zooepidemicus		Yersinia rucken
	Streptococcus equinus		Streptococcus zooepidemicus (Group		Yersinia species
55			_ C)		Yokenella regensburgei
	nonenterococci)	113		175	Yokenella regensburgei (Koserella trabulsii)
	Streptococcus equisimilis		Streptomyces species	1/3	Zygoascus hellenicus
	Streptococcus equisimulis (Group		Suttonella (Kingella) indologenes Tatumella ptyseos		Zygosaccharomyces species
	C/Group G Streptococcus)		raturrena pryseus		_, 300200.12.2, 200 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
		,			

The list includes microorganisms that may be identified by API identification test systems and VITEK® automated identification system from bioMérieux Inc., or by the MicroScan® - WalkAway® automated systems from Dade Behring. Identification relies on classical identification methods using batteries of biochemical and other phenotypical tests.

Table 16. tuf gene sequences obtained in our laboratory (Example 42).

Species	Strain no.	Gene	GenBank Accession no.*
Abiotrophia adiacens	ATCC49175	tuf	AF124224
Enterococcus avium	ATCC14025	tufA	AF124220
		tufB	AF274715
Enterococcus casseliflavus	ATCC25788	tufA	AF274716
		tufB	AF274717
Enterococcus cecorum	ATCC43198	tuf	AF274718
Enterococcus columbae	ATCC51263	tuf	AF274719
Enterococcus dispar	ATCC51266	tufA	AF274720
		tufB	AF274721
Enterococcus durans	ATCC19432	tufA	AF274722
		tufB	AF274723
Enterococcus faecalis	ATCC29212	tuf	AF124221
Enterococcus faecium	ATCC 19434	tufA	AF124222
		tufB	AF274724
Enterococcus gallinarum	ATCC49573	tufA	AF124223
ga		tufB	AF274725
Enterococcus hirae	ATCC8043	tufA	AF274726
Z.//.o/.0000000		tufB	AF274727
Enterococcus malodoratus	ATCC43197	tufA	AF274728
20.0000000		tufB	AF274729
Enterococcus mundtii	ATCC43186	tufA	AF274730
2.110,0000000		tufB	AF274731
Enterococcus pseudoavium	ATCC49372	tufA	AF274732
Emorosous positives		tufB	AF274733
Enterococcus raffinosus	ATCC49427	tufA	AF274734
Zmoroboda ramii ada		tufB	AF274735
Enterococcus saccharolyticus	ATCC43076	tuf	AF274736
Enterococcus solitarius	ATCC49428	tuf	AF274737
Enterococcus sulfureus	ATCC49903	tuf	AF274738
Lactococcus lactis	ATCC11154	tuf	AF274745
Listeria monocytogenes	ATCC15313	tuf	AF274746
Listeria seeligeri	ATCC35967	tuf	AF274747
Staphylococcus aureus	ATCC25923	tuf	AF274739
Staphylococcus epidermidis	ATCC14990	tuf	AF274740
Streptococcus mutans	ATCC25175	tuf	AF274741
Streptococcus pneumoniae	ATCC6303	tuf	AF274742
Streptococcus pyogenes	ATCC19615	tuf	AF274743
Streptococcus suis	ATCC43765	tuf	AF274744

^{*}Corresponding sequence ID NO. for the above ATCC strains are given in table 7.

Table 17. tuf g ne sequenc s selected from databases for Exampl 42.

Speci s	Gene	Accession no.
robacterium tumefaciens	tufA	X99673
•	tufB	X99674
nacystis nidulans	tuf	X17442
Aguifex aeolicus	tufA	AE000657
	tufB	AE000657
Bacillus stearothermophilus	tuf	AJ000260
Bacillus subtilis	tuf	AL009126
Bacteroides fragilis	tuf	P33165
Borrelia burgdorferi	tuf	AE000783
Brevibacterium linens	tuf -	X76863
Bulkholderia cepacia	tuf	P33167
Campylobacter jejuni	tufB	Y17167
	tuf	AE001363
Chlamydia pneumoniae	tuf	M74221
Chlamydia trachomatis	tuf	X77034
Corynebacterium glutamicum	tuf	X77035
Cytophaga lytica	tuf	·AE000513
Deinococcus radiodurans	tui tufA	J01690
Escherichia coli		J01717
	tufB	
Fervidobacterium islandicum	tuf ***fA	Y15788
Haemophilus influenzae	tufA	L42023
	tufB	L42023
Helicobacter pylori	tuf	AE000511
Homo sapiens (Human)	EF-1α	X03558
Methanococcus jannaschii	EF-1α	U67486
Mycobacterium leprae	tuf	D13869
Mycobacterium tuberculosis	tuf	X63539
Mycoplasma genitalium	tuf	L43967
Mycoplasma pneumoniae	tuf	U00089
Neisseria gonorrhoeae	tufA	L36380
Nicotiana tabacum (Tobacco)	EF-1α	U04632
Peptococcus niger	tuf	X76869
Planobispora rosea	. tuf1	U67308
Saccharomyces cerevisiae (Yeast)	EF-1α	X00779
Salmonella typhimurium	tufA	X55116
Jannon Jan Springer	tufB	X55117
Shewanella putrefaciens	tuf	P33169
Spirochaeta aurantia	tuf	X76874
Spirotriaeta aurartua Spirulina platensis	tufA	X15646
Streptomyces aureofaciens	tuf1	AF007125
Streptomyces cinnamoneus	tuf1	X98831
Streptomyces coelicolor	tuf 1	X77039
suapidinycas coalicului	tuf3	X77040
Strontomyoes collinus	tuf1	S79408
Streptomyces collinus	tuf1	X67057
Streptomyces ramocissimus		X67058
	tut2	X67059
O the security and	tuf3	AB001339
Synechocystis sp.	tuf ****f	X77036
Taxeobacter ocellatus	tuf * f	
Thermotoga maritima	tuf	AE000512
Thermus aquaticus	tuf	X66322
Thermus thermophilus	tuf	X06657
Thiobacillus cuprinus	tuf	U78300
Treponema pallidum	tuf	AE000520
Wolinella succinogenes	tuf	X76872

Sequence data were obtained from GenBank, EMBL, and SWISSPROT databases. Genes were designated as appeared in the references.

Tabl 18. Nucl tide and amino acid s quence identities of EF-Tu b tween different nt rococci and other I w G+C gram-positive bact ria.

The upper right triangle represents the deduced amino acid sequence identities of gram-positive bacterial EF-Tu, while the lower left triangle represents the DNA sequence identities of the corresponding tuf genes. The sequence identities between different enterococcal tufA genes are boxed while those between enterococcal tufB genes are shaded.

Becterist tul gene	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15 16	17	18	19	20	21	22	23	24	25	26	27	28	29	30	31	322	33 :	4 3	3 36	37	31	3 31
1. E. avium tudA		96	98	96	96	96	96	97	95	98	99	95			94 96								86			87	86	92	91	90	90	90 8				_	
2. E. cassetflavus IulA	90		87	96	96	89	96	95	96	96	96	95	95		96 94		87				87		87	87	87	88	88	94	91	90	91		2 8				
3. E. dispar rulA	93	90		95	95	96	95	96	95	97	97	91			95 9								87	86	87	87	87	83	80	89	90		2 8		_		
4. E. durans tutA	90	89	90		98	96	99	93	99	95	96	80	91			92							86		87	88	87	94	90	90	90		91 8				
6. E. taecium tulA	89	90	89	96		96	98	93	88		96	69	91	88		92							86		87	88	87	94	85	91	91		33 8				
6. E. galbnarum tulA	90	97	69	89	89		96	83	95	96	96	68	89			82					87		86		87	88	87	93	92	90	90		33 8			-	-
7. E. hirao tulA	90	90	89	99	96	89		93	99	95	96	91	91	89	95 9					86	86		86		87	87	87	94	90	90	90		91 8				
8. E. malodoratus tulA	96	91	94	90	89	90	89		92	97		89	89	90	93 9								85	83	85	86	86	92	90	88	88		91 8			_	
9. E. mundiii tulA	89	89	88	96	93	69	96	88		94	95	88	90	88		1 92				86	96		86		87	88	87	94	80	89	80		91 8				
10.E. pseudozvium tylA	97	92	93	90	89	91	89	97	89		98	90	90	91	95 94					87	87		87	86	87	88	88	93	90	89	90	•	91 B	-			
11.E. rattinosus tutA	97	91	93	90	89	89	89	97	88	97		91	90	90	94 9	93	86			86	86		88		87	87	87	93	89	89	90		91 B			8	
12.E. cecorum tulA	90	90	95	96	96	95	96	92	95	85	95		98	95	93 9					87	87		86	89	87	69	89	93	90	90	91		93 8				
13.E. columbae tidA	90	90	95	96	97	96	96	93	95	95	95	97		95	94 9			88	88	87	88		87	87	87	89	89	94	92	91	91		33 B			_	
14.E. Inocalis tulA	91	91	80	89	98	97	94	94	94	95	96	90	89		94 9	4 j 93	87	87	86	87	27	86	86	87	87	88	87	93	91	89	90		93 B	-	-	-	
15.E. seccharolyticus NAA	91	91	91	90	87	90	89	91	69	92	91	89	89	92	9			87	85	87	86	84	86		87	87	87	92	90	89	69			4 8		8	_
16.E. sullurous tulA	91	89	90	91	88	88	90	91	89	92	91	88	89	91	94	_J91		84	B١	64	85		84		84	85	85	91	90	87	88		_	2 8		_	
17.E. solsarius tud	83	84	83	83	84	83	82	84	83	84	84	84	83	B4	83 8	3				87	87		87	88	88	88	89	92	91	89	80			6 8			
18.E. avam tufB	77	77	78	78	76	77	78	78	77	78	77	76	78	78	77 7	6 77	: ' :	93		94	94		92	88	93	.68	.97	87	86	87	86			9 81			
19.E. casselflavus tufB	71	72	72	72	70	72	72	70	71	72	72	72	70	72	72 6	8 72	79	•	93	95	95	96	95	93	95	84	94	87	86	88	88			10 SH			
20.E. dispar tufB	76	78	77	77	77	77	77	76	77	76	77	77	77	77	78 7	5 78	.82	79	٠. ٠٠	91,	-01		91	94	92	93	.83	86	B3	85	85			9 8			
21.E. durans N/B	77	78	78	78	78	77	78	77	78	77	78	77	77	78	78 7	5 75	83	80	82	. *	88		. 97	94	97	95	. 94	87	86	88	88			ю 9			
22.E. taecum tufB	76	75				77	76	76	76	75	76	77	77	77	76 7	4 74	80	78	79	86	•	.96	97	95	97	85	94	87	87	88	88			10 9			
23.E. gatinerum tufB	72	73	72			74	72	71	72	72	72	72	72	73	73 7	2 72	78	B1	77 -	81	· 82	• '	94	94	95	85	94	85	87	69	89			10 8			
24.E. hirae tulB	75	74					75	75	76	75	75	74	74	74	75 7	2 74	-80	79	79	84	83	79		93	87	. 63	94	87	85	86	88			9			
25.E. maiodoratus tufB	76	76				77	77	74	77	76	76	77	75	77	77 7	3 78	90	79	63	81	80	77	.79		93	88	97	87	86	87	87	85					
26.E. mundtri tu/B	74	74				74	74	74	74	74	74	74	74	75	74 7	1 73	.80	60	78	85	85	80	84	60		94	94	87	88	88	88	64	86 S	10 9			
27.E. pseudoevrum tuli	77		79			78	77	77	76	78	78	77	77	78	78 7	7 78	91	80	85	84	81	79	80	91	80		96	88	87	88	87	65	87 8	XX B			
28.E. rattinosus tufB	7A	79				77		78	77	79	79	78	78	78	79 7	7 79	:90	79	84	84	81	77.	. 80	90	81	92		87	65		68	64		ю 8			
29.A. actacens tut	88	87				86	86	89	86	88	88	87	88	88	88 9	0 62	77	70	76	77	76	71	73	77	73	78	78		90	88	89	90	91 E	35 B	68	4 B	5 6
30.B. subutis tuf	81							79	79	80	81	80	81	81	80 7	B 78	73	69	73	73	71	70	71	72	71	74	74	78		91	92	90	90 E	32 8	28	3 В	2 8
31.L. monocytogenes tuf	82		82					81	81	81	82	81	81	81	81 7	9 79	76	71	75	75	75	73	74	75	73	78	76	79	82		99	88		34 8			4 8
32.L. socigen tuf	1 az	81						81	82	81	82	81	82	80	81 7	9 79	75	71	76	75	74	73	75	75	73	77	76	79	82	99		88	91 8	34 8	58	58	4 8
33.S. aureus tul	84	84						82	84	83	84	86	88	84	82 8	1 79		69	75	75	73	69	72	74	72	74	74	83	79	81	81		96 8	31 8	28	28	10 B
34.S. epidermidis tul	83	85						82	84	83	83	86	87	85		2 79		69	75	75	73	68	72	74	72	74	75	81	79	82	81	94		33 B			
35.S. mutans tuf	76	77				-		75	76	76	76	77	76	76		4 78		72	77	78	77	74	75	78	75	78	81	77	75	78	77	74	73	9	7 9		
36.S. pneumonies tul	76	77				77	77	75	78	76	76	77	76	77		4 75		72	76	78	76	73	74	77	75	75	78	75	76	77	76	74	74 8	37	9	6 9	6 8
	78	77						74	77	76	75	76	75	"		3 79			75	78	75	73	74	75	75		77	76			76	73	72 8	37 9	3	9	M 8
37.S. pyogenes tul	74	78				-		74	76	76	77	77	75	78		3 7			75	78	74	70	74	75	73	73	77	77	77	77	77	72	73 8	38 9	3 9	1	8
38.S. aus tul		76			75							77			75 7										75			75		77	76		74	0 0	3 8	2 0	

Table 19. Strains analyzed in Example 43.

Taxon	Strain*	Strain†	16S rDNA sequence accessi n number
Cedecea	ATCC 33431 ^T		
Cedecea lapagei	ATCC 33432 ^T		
Cedecea neteri	ATCC 33855 ^T	_	
Citrobacter amalonaticus	ATCC 25405 ^T	CDC 9020-77 ^T	AF025370
Citrobacter braakii	ATCC 43162	_	
		CDC 080-58 ^T	AF025368
Citrobacter farmeri	ATCC 51112 ^T	CDC 2991-81 ^T	AF025371
Citrobacter freundii	ATCC 8090 [™]	DSM 30039 ^T	AJ233408
Citrobacter koseri	ATCC 27156 ^T	_	
Citrobacter sedlakii	ATCC 51115 ^T	CDC 4696-86 ^T	AF025364
Citrobacter werkmanii	ATCC 51114 ^T	CDC 0876-58 ^T	AF025373
Citrobacter youngae	ATCC 29935 ^T		
Edwardsiella hoshinae	ATCC 33379 ^T		
Edwardsiella tarda	ATCC 15947 ^T		
	_	CDC 4411-68	AF015259
Enterobacter aerogenes	ATCC 13048 ^T	JCM 1235 [™]	AB004750
Enterobacter agglomerans	ATCC 27989_		
Enterobacter amnigenus	ATCC 33072 ^T	JCM 1237 ^T	AB004749
Enterobacter asburiae	ATCC 35953 ^T	JCM 6051 [™]	AB004744
Enterobacter cancerogenus	ATCC 35317 ^T		
Enterobacter cloacae	ATCC 13047 ^T		
Enterobacter gergoviae	ATCC 33028 ^T	JСМ 1234 ^Т	AB004748
Enterobacter hormaechei	ATCC 49162 ^T	-	
Enterobacter sakazakii	ATCC 29544 ^T	JCM 1233 ^Т	AB004746
Escherichia coli	ATCC 11775 ^T	ATCC 11775 ^T	X80725
Escherichia coli	ATCC 25922	ATCC 25922	X80724
Escherichia coli (ETEC)	ATCC 35401		
Escherichia coli (O157:H7)	ATCC 43895	ATCC 43895	Z83205
Escherichia fergusonii	ATCC 35469 ^T		
Escherichia hermanii	ATCC 33650 ^T		,
Escherichia vulneris	ATCC 33821 ^T	ATCC 33821 ^T	X80734
Ewingella americana	ATCC 33852 ^T		,
	-	NCPPB 3905	X88848
Hafnia alvei	ATCC 13337 ^T	ATCC 13337 ^T	M59155
Klebsiella omithinolytica	ATCC 31898	<u> </u>	
		CIP 103.364	U78182
Klebsiella oxytoca	ATCC 33496	ATCC 13182 ^T	
	-		U78183
Klebsiella planticola	ATCC 33531 ^T	JCM 7251 ^T	AB004755
Klebsiella pneumoniae			
subsp. pneumoniae	ATCC 13883 ^T	DSM 30104 ^T	AJ233420
subsp. ozaenae	ATCC 11296 ^T	ATCC 11296 ^T	Y17654
subsp. rhinoscleromatis	ATCC 13884 ^T		

Tabl 19. Strains analyz d in Exampl 43 (continued).

Taxon	Strain*	Strain†	16S rDNA s quence accessi n number
Kluyvera ascorbata	ATCC 33433 ^T		
		ATCC 14236	Y07650
Kluyvera cryocrescens	ATCC 33435 ^T		
Kluyvera georgiana	ATCC 51603 ^T		
Leclercia adecarboxylata	ATCC 23216 ^T		
Leminorella grimontii	ATCC 33999 ^T	DSM 5078 ^T	AJ233421
Moellerella wisconsensis	ATCC 35017 ^T		
Morganella morganii	ATCC 25830 ^T		
Pantoea agglomerans	ATCC 27155 ^T	DSM 3493 ^T	AJ233423
Pantoea dispersa	ATCC 14589 ^T		
Plesiomonas shigelloïdes	ATCC 14029 ^T		
Pragia fontium	ATCC 49100 ^T	DSM 5563 ^T	AJ233424
Proteus mirabilis	ATCC 25933		
Proteus penneri	ATCC 33519 ^T		
Proteus vulgaris	ATCC 13315 ^T	DSM 30118 ^T	AJ233425
Providencia alcalifaciens	ATCC 9886 ^T		
Providencia rettgeri	ATCC 9250		
Providencia rustigianii	ATCC 33673 ^T		
Providencia stuartii	ATCC 33672		
Rahnella aquatilis	ATCC 33071 ^T	DSM 4594 ^T	AJ233426
Salmonella choleraesuis			
subsp. arizonae	ATCC 13314 ^T		
subsp. choleraesuis			
serotype Choleraesuis	ATCC 7001		
serotype Enteritidis‡	ATCC 13076 ^T		
		SE22	SE22
serotype Gallinarum	ATCC 9184		
serotype Heidelberg	ATCC 8326		
serotype Paratyphi A	ATCC 9150		
serotype Paratyphi B	ATCC 8759		
serotype Typhi‡	ATCC 10749		
		St111	U88545
serotype Typhimurium‡	ATCC 14028		
serotype Virchow	ATCC 51955		
subsp. diarizonae	ATCC 43973 [™]		
subsp. houtenae	ATCC 43974 ^T		
subsp. indica	ATCC 43976 ^T		
subsp. salamae	ATCC 43972 ^T		
Serratia fonticola	DSM 4576 ^T	DSM 4576 ^T	AJ233429
Serratia grimesii	ATCC 14460 ^T	DSM 30063 ^T	AJ233430
Serratia liquefaciens	ATCC 27592 ^T		
Serratia marcescens	ATCC 13880 ^T	DSM 30121 ^T	AJ233431
Serratia odorifera	ATCC 33077 ^T	DSM 4582 ^T	AJ233432
Serratia plymuthica	DSM 4540 ^T	DSM 4540 ^T	AJ233433
Serratia rubidaea	DSM 4480 ^T	DSM 4480 ^T	AJ233436
Shigella boydii	ATCC 9207	ATCC 9207	X96965
Shigella dysenteriae	ATCC 11835		
		ATCC 13313 ^T	X96966
		ATCC 25931	X96964

PCT/CA00/01150 WO 01/23604

Tabl 19. Strains analyz d in Exampl 43 (c ntinued).

Taxon	Strain*	Strain†	16S rDNA sequence acc ssion number
Shigella flexneri	ATCC 12022	ATCC 12022	X96963
Shigella sonnei	ATCC 29930 ^T		·
Tatumella ptyseos	ATCC 33301 ^T	DSM 5000 ^T	AJ233437
Trabulsiella guamensis	ATCC 49490 ^T		
Yersinia enterocolitica	ATCC 9610 ^T	ATCC 9610 ^T	M59292
Yersinia frederiksenii	ATCC 33641 ^T		
Yersinia intermedia	ATCC 29909 ^T		
Yersinia pestis	RRB KIMD27		
•		ATCC 19428 ^T	X75274
Yersinia pseudotuberculosis	ATCC 29833 ^T		
Yersinia rohdei	ATCC 43380 ^T	ER-2935 ^T	X75276
Shewanella putrefaciens	ATCC 8071 ^T		
Vibrio cholerae	ATCC 25870		
		ATCC.14035 ^T	X74695

5

T Type strain

*Strains used in this study for sequencing of partial tuf and atpD genes. SEQ ID NOs. for tuf and atpD sequences corresponding to the above reference strains are given in table 7.

†Strains used in other studies for sequencing of 16S rDNA gene. When both strain numbers are on the same row, both

strains are considered to be the same although strain numbers may be different. ‡Phylogenetic serotypes considered species by the Bacteriological Code (1990 Revision).

Table 20. PCR primer pairs used in this study

Prim r	Sequence	Nucl otide positions*	Amplic n length (bp)
SEQ ID NO.			
<i>tuf</i> 664	5'-AAYATGATIACIGGIGCIGCICARATGGA-	271-299	884
697	3' 5'-CCIACIGTICKICCRCCYTCRCG-3'	1132-1156	
atpD			
568	5'-RTIATIGGIGCIGTIRTIGAYGT-3'	25-47	884
567	5'-TCRTCIGCIGGIACRTAIAYIGCYTG-3'	883-908	
700	5'-TIRTIGAYGTCGARTTCCCTCARG-3'	38-61	871
567	5'-TCRTCIGCIGGIACRTAIAYIGCYTG-3'	883-908	

^{*}The nucleotide positions given are for *E. coli tuf* and *atpD* sequences (GenBank accession no. AE000410 and V00267, respectively). Numbering starts from the first base of the initiation codon.

Table 21. Selection f M. catarrhalis-specific primer pairs from SEQ ID NO: 29¹ (466 pb DNA fragment) other than those previously tested².

		-	-	\mid	-	\vdash	-		L					
Primer	Sequence	Amplicon size (bp)	Moraxella catarrhalis	Moraxella catarrhalis ATCC 53879	Moraxella nonliquefaciens Moraxella lacunata	Moraxella osloensis	Moraxella atlantae	Moraxella phenyipyruvica	Kingella indologenes	Kingella kingea	sibiîigninəm shəssiəM	Asseria gonormosas	Escherichia coll	Staphylococcus aureus
SEO ID NO:118	CGCTGACGGCTTGTTGTACCA	110	6,	+	-			•		•			•	•
SEO ID NO: 119	TGTTTTGAGCTTTTTTTTTTTGA	01-	-		\dashv	+	+	+	1	\downarrow	1	1		
VBmcat1	TGCTTAAGATTCACTCTGCCATTTT	8	+	+	•		<u> </u>		•	•	•		•	•
VBmcat2	TAAGTCGCTGACGGCTTGTTT	S			+	+	+	\downarrow	-	1	\perp	1		
VBmcat3	CCTGCACCACAGTCATCAT	140	+	+	•		<u> </u>			<u>'</u>	•	• •	•	•
VBmcat4	AATTCACCAACAATGTCAAAGC	2			\dagger	+	+	+	+	1		1		
VBmcat5	AATGATAACCAGTCAAGCAAGC	210	+	+			<u> </u>		<u>.</u>	<u>.</u>	<u>'</u>	•	•	٠
VBmcat6	GGTGCATGGTGATTTGTAAAA	2			\dagger	+	+	+	\downarrow	\downarrow	\downarrow	\downarrow		
VBmcat7	GTGTGCGTTCACTTTTACAAAT	160	+	+	•		<u> </u>			•	•	•	•	•
VBmcat8	GGTGTTAAGCTGATGATGAGAG	3			\dagger	\dagger	+	╁	\downarrow	-	1	\downarrow		
VBmcat9	TGACCATGCACCCTTATT	167	+	+			.		<u> </u>		•	•	•	
VBmcat10	TCATTGGGATGAAAGTATCGTT				1	1	-	\dashv	$\frac{1}{2}$	1				

¹ SEQ ID NO. from US patent 6,001,564.

² All PCR assays were performed with 1 ng of purified genomic DNA by using an annealing temperature of 55°C and 30 cycles of amplification. The genomic DNA from the various bacterial species above was always isolated from reference strains obtained from ATCC.

³ All positive results showed a strong amplification signal with genomic DNA from the target species M. catarrhalis.

Table 22. Selection of S. epidermidis-specific primer pairs from SEQ ID NO: 36¹ (705 pb DNA fragment) other than those previously tested.

			1	_1				_						
(O°) ⁵ enutereques toilseannA		द्ध	23	8	22	8	55	8	92	55	9	92	55	
Streptococcus pyogenes		•	<u> </u>	-	<u> </u>	퇴		보	Ŋ		Z	N	•	
Streptococcus pneumoniae		-		-	<u> </u>	틸		F	Ę	•	T N	Ž	•	
Streptococcus agalactiae		•	-			퇸	-	N.	I NT	•	I NT	Ž	•	
Listeria monocytogenes		-	-	-	•	틸	•	TNT	Z	•	N	Z		
Enterococcus gallinarum		-		-	•	Σ	•	TNT	N		TN	<u>Z</u>		
Enterococcus faecalls Enterococcus faecium		: 	<u> </u>	\dashv	-	N N	•	TN TN	NT NT	•	NT NT	N N N		
Bacillus subtilis		$\dot{\exists}$	$\stackrel{\cdot}{\dashv}$	\exists	•	Z Ż	•	Z L N	Z L	•	Z	본	<u>·</u>	
Staphylococcus warner			•	\exists	•	뒫	•	TN TN	FN FN	•	FN FN	Ę	-	
Staphylococcus simulans		-	-		•	-	+	+	-		-	•		
Staphylococcus		\exists		-	•		•	•	•	+	•		,	
Staphylococcus			•	$\overline{\cdot}$	•		•			+	+			
Staphylococcus hominis			•	7	+	+	•	Ę	Ę	•	눌	Ę		
Staphylococcus		•	+	•	+	+	+	•	•	+	+	•	•	
Staphylococcus auricularis		•	•	•	•	•	•	눌	눌	•	Ę	Z	•	
Staphylococcus aureus		•	•	•	+	+	•	•	•	•	•	•	•	
Staphylococcus cohnli		•	٠	•	+	+	-	Ę	Ę	·	Z	NT	•	
Staphylococcus capitls		•	٠	•	+	+	Ŀ	뉟	Z	•	둗	본	٠	
Staphylococcus epidermidis		+	+	+	+	+	+	+	+	+	+	+	+	•
Staphylococcus epidermidis	•	°+	+	+	+	+	+	+	+	+	+	+	+	-
Amplicon size (bp)		125	000	200	900	202		177			153		135	3
Sequence (all 25 nucleotides)	ATCAAAAAGTTGGCGAACCTTTTCA	CAAAAGAGCGTGGAGAAAAGTATCA	CATAGTCTGATTGCTCAAAGTCTTG	GCGAATAGTGAACTACATTCTGTTG	CACGCTCTTTTGCAATTTCCATTGA	GAAGCAAATATTCAAAATGCACCAG	AAAGTCTTTTGCTTCAGATTCA			GAGCATCCATACCTGTGAACACAGA	正していませんべいないない。	1110044114044640410401	TTTGAATTCGCATGTACTTTG	CCCCGGGTTCGAAATCGATAAAAAG
Primer	SEQ ID NO:145	SEQ ID NO:146	VBsep3	VBsep4	VBsep5	VBsep6	VBsep7	VBcon8	odeso.	VBsep9	,	Vasepio	VBsep11	VBsep12

SEQ ID NO. from US patent 6,001,564.

NT = not tested.

² All PCR assays were performed with 1 ng of purified genomic DNA by using an annealing temperature of 55 to 65°C and 30 cycles of amplification. The genomic DNA from the various bacterial species above was always isolated from reference strains obtained from ATCC.

³ All positive results showed a strong amplification signal with genomic DNA from the target species S. epidermidis. The instensity of the positive amplification signal with species other than S. epidermidis was variable.

Table 23.

Influence of nucleotide variation(s) on the efficiency of the PCR amplification: Example with SEQ ID NO: 146 from S. epidermidis.

ยกเษกล 20°C Staphylococcus 9,0 Staphyloccus epidermidis² ATCC 14990 55°C \$ 0, 4 4 4 4 4 \$ 4 2 4 4 4 \$ 4 **ф** ÷ # # # ф , ф \$ \$ # ಕ್ಕ ф ф **20°C** 3+ န 3+4 ŧ ಕ , ŧ **#** # # ф # \$ ŧ, Number of mutation 0 0 2 CAAAGGAGCGTGGTGAAAAGTAGCA CAAAGGAGCGCGGAGAGAAGTACCA CAAAGGAGCGGGAGAAAAGTATCA CAAAAGAGCGTGGAGGAAGTACCA CAAAAGAGCGCGGAGAGAAGTATCA CAAAAGAGCGTGGAGAAAAATATCA CAAAAGAGCGTGGAGAGAAGTATCA CAAAAGAGCGTGGTGAAAAGTATCA CAAAAGAACGTGGAGAAAAGTATCA CAAAGGAGCGTGGAGAAAAGTATCA CHAAAGAGCGTGGAGAAAAGTATCA ATCAAAAGTTGGCGAACCTTTTCA CAAAAGAGCGTGGAGAAAAAGTATCA CAAAAGGGGTGGAGAAAAGTACCA CAAAAGAGCGCGGGAGAAAGTATCA Sequence (all 25 nucleotides) **SEQ ID NO:145 SEQ ID NO:146** VBmut12 VBmut13 VBmut 10 VBmut11 VBmut8 VBmut2 VBmut3 VBmut4 VBmut5 **VBmut6** VBmut9 VBmut1 VBmut7 Primer¹

All PCR tests were performed with SEQ ID NO:145 without modification combined with SEQ ID NO:146 or 13 modified versions of SEQ ID NO:146. Boxed nucleotides indicate changes in SEQ ID NO:146. All SEQ ID NOs. are from US patent 6,001,564 The tests with S. epidermidis were performed by using an annealing temperature of 55°C with 1, 0,1 and 0,01 ng of puritied genomic DNA or at 50°C with 1 ng of puritied genomic DNA.

³ The tests with S. aureus were performed only at 50°C with 1 ng of genomic DNA.

4 The intensity of the positive amplification signal was quantified as follows: 3+ = strong signal, 2+ = intermediate signal and + = weak signal.

Effect of the primer length on the efficiency of the PCR amplification¹: Example with the AT-rich SEQ ID NO: 145² and SEQ ID NO: 146² from S. epidermidis. Table 24.

				Sta e ₉ A	phylo sidem TCC 1	Staphylococcus epidermidis³ ATCC 14990	to.		hylococcus aureus		ουγίος γαθετίς της της της της της της της της της τη	υμλιοcoccns csbirls		phylococcus warnen	
		1 enath		45°C			55°C		de1S		dejs	1612	Ima	Stap	
Primer	Sequence	(nt)	-	1,0	0,01	-	0,1	0,01	45 5	55 45	52	45	22	45	55
VBsep301	ATATCATCAAAAAGTTGGCGAACCTTTTCA	98	!	!	!	 		-	Ļ	:	_	!		ţ	
VBsep302	AATTGCAAAAGAGCGTGGAGAAAAGTATCA	30	Ż	Z	Ē	++	3+	5+	2	Ž	•	ž	•	ž	
SEQ ID NO:145	ATCAAAAGTTGGCGAACCTTTTCA	25			,										
SEQ ID NO:146	CAAAAGAGCGTGGAGAAAAGTATCA	25	4	*	5 +	‡	÷	5 +	•		•	+	•	•	
VBsep201	AAAGTTGGCGAACCTTTTCA	20	!	!	!	-			•	:		ŀ		Ė	
VBsep202	GAGCGTGGAGAAAGTATCA	20	Z	Z	Z	++	÷	+7	Ž	Z	•	ž		Ē	
VBsep171	GTTGGCGAACCTTTTCA	17		•	,										
VBsep172	CGTGGAGAAAGTATCA	17	+	3+	5 +	÷	+7	+			•		•		
VBsep151	TGGCGAACCTTTTCA	15		•											
VRsen152	TGGAGAAAGTATCA	15	,	<u>+</u>	+	•				<u>. </u>		•		•	•

¹ All PCR tests were performed using an annealing temperature of 45 or 55°C and 30 cycles of amplification.

NT = not tested.

² All SEQ ID NOs. in this Table are from US patent 6,001,546.

³ The tests with S. epidemidis were made with 1, 0,1 and 0,01 ng of purified genomic DNA.

⁴ The tests with all other bacterial species were made only with 1 ng of purified genomic DNA.

⁵ The Intensity of the positive amplification signal was quantified as follows: 4+ = very strong signal, 3+ = strong signal, 2+ = intermediate signal and + = weak signal.

Effect of the primer length on the efficiency of the PCR amplification[†]: Example with the GC-rich SEQ ID NO: 83² and SEQ ID NO: 84² from P. aeruginosa. Table 25.

	T				7		
ophilus parahaemolyticus						•	
eibiligninəm site			\dashv	•	-	•	\dashv
silinqotlam esnomondot			-		_		\dashv
ebitug silan	SWƏYS				'		
olderia cepacia			_		-		
omonas fluorescens ⁴	Psend				_		_
onas sa 554	0,01		•		•	,	·
Pseudomonas aeruginosa ATCC 35554	ATCC 35554 1 0,1 0,0			•	+	•	F
				Ġ	* >	ć	<u> </u>
	Length (nt)	19	19	16	16	13	13
	Sequence	CGAGCGGTGTGTTCATC	CAAGTCGTCGTCGGAGGGA	CGAGCGGCTGCTTC	GTCGTCGCAGGGA	GCGGGTGTTC	GTCGTCGGAGGGA
	Primer	SEQ ID NO 83	SEQ ID NO 84	Pse554-16a	Pse674-16a	Pse554-13b	Pse674-13a

¹ All PCR tests were performed using an annealing temperature of 55°C and 30 cycles of amplification.

² All SEQ ID NOs. in this Table are from US patent 6,001,546.

³ The tests with P. aeruginosa were made with 1, 0,1 and 0,01 ng of purified genomic DNA.

4 The tests with all other bacterial species were made only with 1 ng of purified genomic DNA.

⁵ The intensity of the positive amplification signal was quantified as follows: 2+ = strong signal and + = moderately strong signal.

Annex I: Specific and ubiquitous primers for nucleic acid amplification (tuf sequences).

			Originating DNA fragment
5	SEQ ID NO.	Nucleotide sequence	SEQ ID Nucleotide NO. position
0	Bacterial s	species: Acinetobacter bauman	nii
	1692	5'-GGT GAG AAC TGT GGT ATC TTA CTT	1 478-501
	1693 ^a	5'-CAT TTC AAC GCC TTC TTT CAA CTG	1 691-714
5	Bacterial s	species: Chlamydia pneumoniae)
		5'-CGG AGC TAT CCT AGT CGT TTC A	20 2-23
	630 629 ^a	5'-AAG TTC CAT CTC AAC AAG GTC AAT A	
			20 45-68
20	2085	5'-CAA ACT AAA GAA CAT ATC TTG CTA 5'-ATA TAA TTT GCA TCA CCT TCA AG	20 237-259
	2086 ^a	5'-ATA TAA TIT GCA TCA CCT TCA NO	
	2087	5'-TCA GCT CGT GGG ATT AGG AGA G	20 431-452 20 584-605
05	2088 ^a	5'-AGG CTT CAC GCT GTT AGG CTG A	20 584-605
25	Bacterial s	species: Chlamydia trachomati	is
	554	5'-GTT CCT TAC ATC GTT GTT TTT CTC	22 82-105
	555 ^a	5'-TCT CGA ACT TTC TCT ATG TAT GCA	22 249-272
30	<u>Parasitica</u>	l species: Cryptosporidium part	vum
	798	5'-TGG TTG TCC CAG CCG ATC GTT T	865 158-179
	804 ^a	5'-CCT GGG ACG GCC TCT GGC AT	865 664-683
35	-	5'-ACC TGT GAA TAC AAG CAA TCT	865 280-300
	799 805 ^a	5'-CTC TTG TCC ATC TTA GCA GT	865 895-914
	805-		865 307-330
	800	5'-GAT GAA ATC TTC AAC GAA GTT GAT	865 307-330 865 946-966
40	806 ^a	5'-AGC ATC ACC AGA CTT GAT AAG	
	801	5'-ACA ACA CCG AGA AGA TCC CA	865 353-372
	803a	5'-ACT TCA GTG GTA ACA CCA GC	865 616-635
45	802	5'-TTG CCA TTT CTG GTT TCG TT	865 377-396
7.5	807 ^a	5'-AAA GTG GCT TCA AAG GTT GC	865 981-1000
	Bacterial	species: Enterococcus faeciu	m
50	1696	5'-ATG TTC CTG TAG TTG CTG GA	64 189-208
J0		5'-TTT CTT CAG CAA TAC CAA CAA C	64 422-443
	Bacterial	species: Klebsiella pneumoni	ae
£ £		5'-TGT AGA GCG CGG TAT CAT CAA AGT	A 103 352-377
55	1329 1330 ^a	5'-AGA TTC GAA CTT GGT GTG CGG G	103 559-571

These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex I: Specific and ubiquitous primers for nucleic acid amplification (tuf sequences) (continued).

				Originating	DNA fragment
5	SEQ ID NO.	Nucleotide	sequence	SEQ ID	Nucleotide position
10	Bacterial	species:	Mycoplasma pneumoniae		
•	2093 2094 ^b		CAA TCG AAG ACA CC	2097 ^a 2097 ^a	635-654 709-732
	2094~	5'-TIC AAT	TTC TTG ACC TAC TTT CAA	2037	705 732
15	<u>Bacterial</u>	species:	Neisseria gonorrhoeae		
	551	5'-GAA GAA	AAA ATC TTC GAA CTG GCT A	126	256-280
	552 ^b	5'-TAC ACG	GCC GGT GAC TAC G	126	378-396
20	2173	5'-AAG AAA	AAA TCT TCG AAC TGG CTA	126	257-280
	2174 ^b	5'-TCT ACA	CGG CCG GTG	126	384-398
	2175	5'-CCG CCA	TAC CCC GTT T	126	654-669
0.	2176 ^b	5'-CGG CAT	TAC CAT TTC CAC ACC TTT	126	736-759
25	Bacterial	species:	Pseudomonas aeruginosa	ı	
	1694		AGG ATG ACA ACG GC	153	231-250
30	1695 ^b	5'-ACG ATT	TCC ACT TCT TCC TGG	153	418-438
30	Bacterial	species:	Streptococcus agalacti	ae	
	549 .	5'-GAA CGT	GAT ACT GAC AAA CCT TTA	207-210 ^C	308-331 ^d
25	550 ^b	5'-GAA GAA	GAA CAC CAA CGT TG	207-210 ^C	520-539 ^d
35	Bacterial	species:	Streptococcus pyogenes	3	
	999	5'-TTG ACC	TTG TTG ATG ACG AAG AG	1002	143-165
40	1000 ^b	5'-TTA GTG	TGT GGG TTG ATT GAA CT	1002	622-644
40	1001	5'-AAG AGT	TGC TTG AAT TAG TTG AG	1002	161-183
	1000 ^b	5'-TTA GTG	TGT GGG TTG ATT GAA CT	1002	622-644
45	<u>Parasitic</u>	al species:	Trypanosoma brucei		
43	820	5'-GAA GGA	GGT GTC TGC TTA CAC	864	513-533
	821 ^b	5'-GGC GCA	AAC GTC ACC ACA TCA	864	789-809
	820	5'-GAA GGA	GGT GTC TGC TTA CAC	864	513-533
50	822 ^b	5'-CGG CGG	ATG TCC TTA ACA GAA	864	909-929

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

c These sequences were aligned to derive the corresponding primer.

d The nucleotide positions refer to the S. agalactiae tuf sequence fragment (SEQ ID NO. 209).

Annex I: Specific and ubiquitous primers for nucleic acid amplification (tuf sequences) (continued).

									Originating I	ONA fragment
	SEQ ID NO.	Nucleotide	sequen	ce					SEQ ID NO.	Nucleotide position
	Parasitical	species:	Tryp	anosc	ma (cruz	i			
	794	5'-GAC GAC	AAG TC	G GTG	AAC	тт			840-842ª	
	795 ^b	5'-ACT TGC	ACG CG	A TGT	GGC	AG			840-842 ^a	874-893 ^C
	Bacterial ge	nus:	Clos	tridi	um :	sp.				
	796	5'-GGT CCA	ATG CC	W CAA	ACW	AGA			32,719- 724,736 ^a	32-52 ^d
)	797b	5'-CAT TAA	GAA TG	G YTT	ATC	TGT	SKC	TCT	32,719- 724,736 ^a	320-346 ^d
	808	5'-GCI TTA	. IWR GC	А ТТА	GAA	RAY	CCA		32,719- 724,736 ^a	224-247 ^d
	809b	5'-TCT TCC	TGT WG	C AAC	TGT	TCC	TCT		32,719- 724,736 ^a	337-360 ^d
5	810	5'-AGA GMW	ACA GA	T AAR	SCA	TTC	TTA		32,719- 724,736 ^a	320-343 ^d
	811 ^b	5'-TRA ART	' AGA AT	T GTG	GTC	TRT	ATC	С	32,719- 724,736 ^a	686-710 ^d
)	Bacterial ge	nus:	Cory	nebad	cter	ium	sp.			
	545	5'-TAC ATO	CTB GT	Y GCI	CTI	AAC	AAG	TG	34-44,662 ^a	
	546 ^b	5'-CCR CG	CCG G1	R ATG	GTG	AAG	AT		34-44,662 ^a	350-372 ^e
5	Bacterial ge	enus:	Ente	rocod	ccus	sp.	•			
	656	5'-AAT TA	TGG CT	G CAG	TTG	AYG	A		58-72ª	273-294 ^f
)	657 ^b	5'-TTG TCC	ACG T	C GAT	RTC	TTC	A		58-72 ^a	556-577 [£]
,	656	5'-AAT TA	TGG CI	G CAG	TTG	AYG	A		58-72 ^a	273-294 ^f
	271 ^b	5'-TTG TCC	ACG T	G GAT	RTC	TTC	A		58-72 ^a	556-577 [£]
	1137	5'-AAT TA	TGG C	rg CWG	TTG	AYG	AA		58-72ª	273-295 ^f
5	1136 ^b	5'-ACT TG	CCA CC	T TSG	ATR	TCT			58-72 ^a	559-579 [£]

a These sequences were aligned to derive the corresponding primer.

55

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

 $^{^{\}rm C}$ The nucleotide positions refer to the T. cruzi tuf sequence fragment (SEQ ID NO. 842).

d The nucleotide positions refer to the *C. perfringens* tuf sequence fragment (SEQ ID NO. 32).

^e The nucleotide positions refer to the C. diphtheriae tuf sequence fragment (SEQ ID NO. 662).

f The nucleotide positions refer to the E. durans tuf sequence fragment (SEQ ID NO. 61).

Annex I: Sp cific and ubiquitous primers for nucleic acid amplification (tuf sequences) (continued).

					Originating	DNA fragment
	SEQ ID NO.	Nucleotide	e sequence		SEQ ID NO.	Nucleotide position
. 10	Bacterial ge	enus:	Legione.	lla sp.		
	2081	5'-GRA TYI	TYA AAG TI	rg gtg agg aag	111-112 ^a	
	2082 ^C	5'-CMA CT	CAT CYC GO	CT TCG TAC C	111-112 ^a	548-569b
15	Bacterial go	enus:	Staphyl	ococcus sp.		
	553	5'-GGC CG	GTT GAA CO	GT GGT CAA ATC A	176-203 ^a	313-337 ^d
	575 ^C	5'-TIA CC	A TTT CAG TA	AC CTT CTG GTA A	176-203 ^a	653-677 ^d
20	553	5'-GGC CG'	r GTT GAA CO	GT GGT CAA ATC A	176-203 ^a	313-337d
20	707 ^C			AC CTT CTG GTA A		653-677 ^d
	Bacterial g	enus:	Strepto	coccus sp.		
25	547	5'-GTA CA	TTG CTT C	AG GAC GTA TC	206-231 ^a	372-394 ^e
	548 ^C	5'-ACG TT	C GAT TTC A	TC ACG TTG	206-231 ^a	548-568 ^e
	Fungal genu	<u>s</u> :	Candida	sp.	•	
30	576	5'-AAC TT	C RTC AAG AA	AG GTY GGT TAC A	A 407-426, 428-432 ^a	332-357 ^f
	632 ^c	5'-CCC TT	r GGT GGR TO	CS TKC TTG GA	407-426, 428-432 ^a	791-813 [£]
35	631	5'-CAG AC	C AAC YGA I	AA RCC ATT RAG A	AT 407-426, 428-432 ^a	523-548 ^f
	632 ^c	5'-CCC TT	T GGT GGR T	CS TKC TTG GA	407-426, 428-432 ^a	791-813 [£]
40	633	5'-CAG AC	C AAC YGA I	AA RCC ITT RAG A	AT 407-426, 428-432 ^a	523-548 [£]
	632 ^c	5'-CCC TT	T GGT GGR T	CS TKC TTG GA	407-426, 428-432 ^a	791-813 ^f

a These sequences were aligned to derive the corresponding primer.

45

50

b The nucleotide positions refer to the L. pneumophila tuf sequence fragment (SEQ ID NO. 112).

C These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

d The nucleotide positions refer to the S. aureus tuf sequence fragment (SEQ ID NO. 179).

 $^{^{}m e}$ The nucleotide positions refer to the *S. agalactiae* tuf sequence fragment (SEQ ID NO. 209).

f The nucleotide positions refer to the C. albicans tuf(EF-1) sequence fragment (SEQ ID NO. 408).

Annex I: Specific and ubiquitous primers for nucleic acid amplification (tuf sequences) (continued).

		Originating DNA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID Nucleotide NO. position
Fungal gen	us: Cryptococcus sp.	
1971	5'-CYG ACT GYG CCA TCC TYA TCA	434,623,1281, 150-170 ^b 1985,1986 ^a
1973 ^C	5'-RAC ACC RGI YTT GGW ITC CTT	434,623,1281, 464-484 ^b 1985,1986 ^a
1972	5'-MGI CAG CTC ATY ITT GCW KSC	434,623,1281, 260-280 ^b 1985,1986 ^a
1973 ^c	5'-RAC ACC RGI YTT GGW ITC CTT	434,623,1281, 464-484 ^b 1985,1986 ^a
<u>Parasitica</u>	l genus: Entamoeba sp.	
703 704 ^C	5'-TAT GGA AAT TCG AAA CAT CT 5'-AGT GCT CCA ATT AAT GTT GG	512 38-57 512 442-461
703 705 ^C	5'-TAT GGA AAT TCG AAA CAT CT 5'-GTA CAG TTC CAA TAC CTG AA	512 38-57 512 534-553
703 706 [©]	5'-TAT GGA AAT TCG AAA CAT CT 5'-TGA AAT CTT CAC ATC CAA CA	512 38-57 512 768-787
793 704 ^C	5'-TTA TTG TTG CTG CTG GTA CT 5'-AGT GCT CCA ATT AAT GTT GG	512 149-168 512 442-461
<u>Parasitica</u>	al genus: Giardia sp.	
816 819 ^C	5'-GCT ACG ACG AGA TCA AGG GC 5'-TCG AGC TTC TGG AGG AAG AG	513 305-324 513 895-914
817 818 ^C	5'-TGG AAG AAG GCC GAG GAG TT 5'-AGC CGG GCT GGA TCT TCT TC	513355-374513825-844
<u>Parasitica</u>	al genus: Leishmania sp.	
701 702 ^C	5'-GTG TTC ACG ATC ATC GAT GCG 5'-CTC TCG ATA TCC GCG AAG CG	514-526 ^a 94-114 ^d 514-526 ^a 913-932 ^d

^{· 50} a These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the *C. neoformans tuf* (EF-1) sequence fragment (SEQ ID NO. 623).

C These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

of the nucleotide positions refer to the L. tropica tuf(EF-1) sequence fragment (SEQ ID NO. 526).

Annex I: Specific and ubiquitous prim rs for nucleic acid amplification (tuf sequences) (continued).

					Originating D	NA fragment
5	SEQ ID NO.	Nucleotide	sequence		SEQ ID NO.	Nucleotide position
10	Parasitical	genus:	Trypanoso	ma sp.		
	823	5'-GAG CGG	TAT GAY GAG	ATT GT	529,840-	493-512 ^b
15	824 ^c		TGC GGC ACC		842,864 ^a 529,840- 842,864 ^a	1171-1190 ^b
	Bacterial f	amily:	Enterobac	teriaceae		
	933	5'-CAT CAT	CGT ITT CMT	GAA CAA RTG	78,103,146, 168,238,698 ^a	390-413 ^d
20	934 ^C	5'-TCA CGY	TTR RTA CCA	CGC AGI AGA	78,103,146, 168,238,698 ^a	831-854 ^d
	Bacterial f	amily:	Mycobacte	riaceae		
25	539	5'-CCI TAC	ATC CTB GTY	GCI CTI AAC AAG	122	85-111
	540 ^C		TCY TCR TCG		122	181-203
	Bacterial c	roup:	Escherich	ia coli and S	higella	
30	1661	5'-TGG GAA	GCG AAA ATC	CTG	1668 ^e	283-300
	1665 ^C		AGG TAG ACT		1668 ^e	484-502
	Bacterial o	group:	Pseudomor	ads group		
35	541	5'-CTK GA	ATG TTC CGC	AAG CTG CT	153-155 ^a	476-498 [£]
	542 ^C		TAG AAC TGS		153-155 ^a	679-702 [£]
		F (CMV CA)	ATG TTC CGC	AAC CTG CT	153-155 ^a	476-498 [£]
40	541 544 ^C		TCG CCM GGC		153-155 ^a	749-771 ^f
70	 -					

a These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the T. brucei tuf (EF-1) sequence fragment (SEQ ID NO. 864).

C These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

d The nucleotide positions refer to the $E.\ coli$ tuf sequence fragment (SEQ ID NO. 698).

e Sequence from databases.

f The nucleotide positions refer to the P. aeruginosa tuf sequence fragment (SEQ ID NO. 153).

Annex I: Specific and ubiquitous primers for nucleic acid amplification (tuf sequences) (continued).

				C	originating D	NA fragment
	SEQ ID NO.	Nucleotide sequence			SEQ ID NO.	Nucleotide position
	Parasitical	group: Trypan	osomatidae	family		
	923	5'-GAC GCI GCC ATC	CTG ATG ATC		511,514-526, 529,840-842, 864 ^a	166-188 ^b
	924 ^C	5'-ACC TCA GTC GTC	ACG TTG GCG		511,514-526, 529,840-842, 864 ^a	648-668 ^b
	925	5'-AAG CAG ATG GTT	GTG TGC TG		511,514-526, 529,840-842, 864 ^a	274-293b
	926 ^c	5'-CAG CTG CTC GTG	GTG CAT CTC G	AT	511,514-526, 529,840-842,	676-699 ^b
;	927	5'-ACG CGG AGA AGG	TGC GCT T		864a 511,514-526, 529,840-842, 864a	
)	928 ^C	5'-GGT CGT TCT TCG	AGT CAC CGC A	4	511,514-526, 529,840-842, 864 ^a	
		Unive	rsal primers	s (bact	eria)	
5	636	5'-ACT GGY GTT GAI	ATG TTC CGY A	AA.	7,54,78, 100,103,159, 209,224,227	
0	· 637 ^C	5'-ACG TCA GTI GTA	CGG AAR TAG A	A.A	7,54,78, 100,103,159 209,224,227	692-714 ⁰
J	638	5'-CCA ATG CCA CAA	ACI CGT GAR (CAC AT	7,54,78, 100,103,159 209,224,227	
5	639 ^c	5'-TTT ACG GAA CAT	TTC WAC ACC	WGT IAC	- · · · · · · · · · · · · · · · · · · ·	469-496

⁵⁰ a These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the *L. tropica tuf* (EF-1) sequence fragment (SEQ ID NO. 526).

C These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

d The nucleotide positions refer to the E. coli tuf sequence fragment (SEQ ID NO. 78).

 $^{^{\}mathrm{e}}$ The nucleotide positions refer to the B. cereus tuf sequence fragment (SEQ ID NO. 7).

Annex I: Specific and ubiquitous prim rs for nucleic acid amplification (tuf sequences) (continued).

			Originating DNA fragment
	SEQ ID NO.	Nucleotide sequence	SEQ ID Nucleotide NO. position
10		Universal primers (bacteria) (c	ontinued)
15	643	5'-ACT GGI GTI GAR ATG TTC CGY AA	1,3,4,7,12, 470-492b 13,16,49,54, 72,78,85,88, 91,94,98,103, 108,112,115, 116,120,121, 126,128,134,
20			136,146,154, 159,179,186, 205,209,212, 224,238 ^a
•	644 ^C	5'-ACG TCI GTI GTI CKG AAR TAG AA	same as SEQ $692-714^{\mathrm{b}}$ ID NO. 643
30	643	5'-ACT GGI GTI GAR ATG TTC CGY AA	1,3,4,7,12, 470-492 ^b 13,16,49,54, 72,78,85,88, 91,94,98,103, 108,112,115, 116,120,121, 126,128,134,
35			136,146,154, 159,179,186, 205,209,212, 224,238 ^a
	645 ^C	5'-ACG TCI GTI GTI CKG AAR TAR AA	same as SEQ 692-714 ^b ID NO. 643
40	646	5'-ATC GAC AAG CCI TTC YTI ATG SC	2,13,82 317-339 ^d 122,145 ^a
	647 ^C	5'-ACG TCC GTS GTR CGG AAG TAG AAC TG	_
45	646	5'-ATC GAC AAG CCI TTC YTI ATG SC	2,13,82 317-339 ^d 122,145 ^a
	648 ^C	5'-ACG TCS GTS GTR CGG AAG TAG AAC TO	3 2,13,82 686-711 ^d 122,145 ^a

a These sequences were aligned to derive the corresponding primer.

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b The nucleotide positions refer to the $E.\ coli\ tuf$ sequence fragment (SEQ ID NO. 78).

C These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

d The nucleotide positions refer to the A. meyeri tuf sequence fragment (SEQ ID NO. 2)

Annex I: Specific and ubiquitous prim rs for nucleic acid amplification (tuf sequences) (continued).

		Originating DNA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID Nucleotide NO. position
	Universal primers (bacteria) (co	ntinued)
649	5'-GTC CTA TGC CTC ARA CWC GIG AGC AC	8,86,141,143 ^a 33-58 ^b
650 ^C	5'-TTA CGG AAC ATY TCA ACA CCI GT	8,86,141,143 ^a 473-495 ^b
636	5'-ACT GGY GTT GAI ATG TTC CGY AA	8,86,141,143 ^a 473-495 ^b
651 ^C	5'-TGA CGA CCA CCI TCY TCY TTY TTC A	8,86,141,143 ^a 639-663 ^b
	Universal primers (fungi)	
1974	5'-ACA AGG GIT GGR MSA AGG AGA C	404,405,433, 443-464 ^d 445,898,1268,
1975 ^C	5'-TGR CCR GGG TGG TTR AGG ACG	1276,1986 ^a 404,405,433, 846-866 ^d 445,898,1268,
		1276,1986 ^a
1976	5'-GAT GGA YTC YGT YAA ITG GGA	407-412, 286-306 ^e 414-426,428- 431,439,443,447, 448,622,624,665,
	•	1685,1987-1990 ^a
1978 ^C	5'-CAT CIT GYA ATG GYA ATC TYA AT	same as SEQ 553-575 ^e ID NO. 1976
1977	5'-GAT GGA YTC YGT YAA RTG GGA	same as SEQ 286-306 ^e ID NO. 1976
1979 ^C	5'-CAT CYT GYA ATG GYA ASC TYA AT	same as SEQ 553-575 ^e ID NO. 1976
1981	5'-TGG ACA CCI SCA AGI GGK CYG	401-405, 281-301 ^d 433,435,436, 438,444,445,449, 453,455,457,779, 781-783,785,786,
		788-790,897-903, 267-1272,1274-1280,
		282-1287,1991-1998 ^a
1980 ^C	5'-TCR ATG GCI TCI AIR AGR GTY T	same as SEQ 488-509 ^d ID NO. 1981

a These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the $B.\ distasonis$ tuf sequence fragment (SEQ ID NO. 8).

⁵⁵ C These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

d The nucleotide positions refer to the A. fumigatus tuf (EF-1) sequence fragment (SEQ ID NO. 404).

e The nucleotide positions refer to the C. albicans tuf (EF-1) sequence fragment (SEQ ID NO. 407).

Annex I: Specific and ubiquitous primers for nucleic acid amplification (tuf sequences) (continued).

SEQ ID NO.	Nucleotide se	quence					SEQ ID NO.	Nucleotic position
	Universal p	rimers	(fung	i)	(cont	inue	i)	_
1982	5'-TGG ACA CY	TI SCA AGI	GGK C	YG			me as SEQ NO. 1981	281-301 ⁶
1980 ^b	5'-TCR ATG GO	TCI AIF	R AGR G	TY ?	r		me as SEQ NO. 1981	488-509
1983	5'-CYG AYT GO	G CYA TIC	TCA T	CA			me as SEQ NO. 1981	143-163
1980 ^b	5'-TCR ATG GO	I TCI AIF	R AGR G	TY T	r		me as SEQ NO. 1981	488-509
1984	5'-CYG AYT GY	G CYA TYO	r, art	rca			ame as SEQ NO. 1981	143-163
1980 ^b	5'-TCR ATG GO	CI TCI AI	R AGR (STY '	Т		ame as SEQ NO. 1981	488-509
	Sequencing	primers	•					
556	5'-CGG CGC N	AT CYT SG	r TGT 7	rgc			668 ^C	306-32
557 ^b	5'-CCM AGG C				G		668 ^C	1047-10
604	5'-CGG CGC I	איזי כעידי פכי	ኮ ጥርጥ '	TGC			668 ^C	306-32
694 557b	5'-CCM AGG C				G		668 ^C	1047-10
664	5'-AAY ATG A	TT ACT GG	T GCI (GCI	CAR A	rg ga	619 ^C	604-63
652 ^b	5'-CCW AYA G						619 ^C	1482-15
664	5'-AAY ATG A	TI ACI GG	I GCI (GCI	CAR A	rg ga	619 ^C	604-63
561b	5'-ACI GTI C						619 ^C	1483-15
543	5'-ATC TTA G	TA GTT TC	T GCT	GCT	GA		607	8-30
660p	5'-GTA GAA T						607	678-70
658	5'-GAT YTA G	TC GAT GA	T GAA	GAA	TT		621	116-13
659 ^b	5'-GCT TTT T	GI GTT TC	w GGT	TTR	ΑT		621	443-46
658	5'-GAT YTA G	TC GAT GA	T GAA	GAA	TT		621	116-13
661 ^b	5'-GTA GAA Y	TG TGG WC	G ATA	RTT	RT		621	678-70
558	5'-TCI TTY A	AR TAY GO	I TGG	GΤ			665 ^C	157-17
559b	5'-CCG ACR G				AT		665 ^C	1279-13
813	5'-AAT CYG T						665 ^C	687-70
559 ^b	5'-CCG ACR G	CR AYI GT	Y TGI	CKC	AT		665 ^C	1279-13

The nucleotide positions refer to the A. fumigatus tuf (EF-1) sequence fragment (SEQ ID NO. 404).

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

C Sequences from databases.

Annex I: Specific and ubiquitous primers for nucleic acid amplification (tuf sequenc s) (continued).

			Originating	DNA fragmen
	SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
		Sequencing primers (continued)		
	558	5'-TCI TTY AAR TAY GCI TGG GT	665 ^a	157-176
	815 ^b	5'-TGG TGC ATY TCK ACR GAC TT	665 ^a	686-705
	560	5'-GAY TTC ATY AAR AAY ATG ATY AC	665 ^a	289-311
	559 ^b	5'-CCG ACR GCR AYI GTY TGI CKC AT	· 665 ^a	1279-1301
	653	5'-GAY TTC ATI AAR AAY ATG AT	665 ^a	289-308
	559b	5'-CCG ACR GCR AYI GTY TGI CKC AT	665 ^a	1279-1301
)	558	5'-TCI TTY AAR TAY GCI TGG GT	665 ^a	157-176
	655b ·	5'-CCR ATA CCI CMR ATY TTG TA	665 ^a	754-773
	654	5'-TAC AAR ATY KGI GGT ATY GG	665 ^a	754-773
5	559b	5'-CCG ACR GCR AYI GTY TGI CKC AT	665 ^a	1279-130
	696	5'-ATI GGI CAY RTI GAY CAY GGI AAR AC	: 698 ^a	52-77
	697 ^b	5'-CCI ACI GTI CKI CCR CCY TCR CG	698 ^a	1132-115
)	911	5'-GAC GGM KKC ATG CCG CAR AC	853	22-41
	914b	5'-GAA RAG CTG CGG RCG RTA GTG	853	700-720
	912	5'-GAC GGC GKC ATG CCG CAR AC	846	20-39
_	914 ^b	5'-GAA RAG CTG CGG RCG RTA GTG	846	692-712
5	913	5'-GAC GGY SYC ATG CCK CAG AC	843	251-270
	915 ^b	5'-AAA CGC CTG AGG RCG GTA GTT	843	905-925
	916	5'-GCC GAG CTG GCC GGC TTC AG	846	422-441
0	561 ^b	5'-ACI GTI CGG CCR CCC TCA CGG AT	619 ^a	1483-150
	664	5'-AAY ATG ATI ACI GGI GCI GCI CAR A'	rg ga 619 ^a	604-632
	917 ^b	5'-TCG TGC TAC CCG TYG CCG CCA T	846	593-614

a Sequences from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex I: Specific and ubiquitous primers for nucleic acid amplification (tuf s quences) (continued).

		Originating DNA fragment
	SEQ ID NO.	Nucleotide sequence SEQ ID Nucleotide NO. position
10		Sequencing primers (continued)
•	1221	5'-GAY ACI CCI GGI CAY GTI GAY TT 1230a 292-314
	1226 ^b	5'-GTI RMR TAI CCR AAC ATY TC 1230 ^a 2014-2033
15	1222	5'-ATY GAY ACI CCI GGI CAY GTI GAY TT 1230ª 289-314
	1223 ^b	5'-AYI TCI ARR TGI ARY TCR CCC ATI CC 1230a 1408-1433
	1224	5'-CCI GYI HTI YTI GAR CCI ATI ATG 1230ª 1858-1881
	1225 ^b	5'-TAI CCR AAC ATY TCI SMI ARI GGI AC 1230ª 2002-2027
20	1227	5'-GTI CCI YTI KCI GAR ATG TTY GGI TA 1230ª 2002-2027
	1229 ^b	5'-TCC ATY TGI GCI GCI CCI GTI ATC AT 698ª 4-29
	1228	5'-GTI CCI YTI KCI GAR ATG TTY GGI TAY GC 1230ª 2002-2030
25	1229 ^b	5'-TCC ATY TGI GCI GCI CCI GTI ATC AT 698ª 4-29
	1999	5'-CAT GTC AAY ATT GGT ACT ATT GGT CAT GT 498-500, 25-53 ^d 502,505,506,
		508,619,2004,2005 ^C
30	2000 ^b	5'-CCA CCY TCI CTC AMG TTG AAR CGT T same as SEQ 1133-1157 ^d ID NO. 1999
	2001	5'-ACY ACI TTR ACI GCY GCY ATY AC same as SEQ 67-89 ^d ID NO. 1999
35	2003 ^b	5'-CAT YTC RAI RTT GTC ACC TGG same as SEQ 1072-1092 ^d ID NO. 1999
	2002	5'-CCI GAR GAR AGA GCI MGW GGT same as SEQ 151-171 ^d ID NO. 1999
40	2003 ^b	5'-CAT YTC RAI RTT GTC ACC TGG same as SEQ 1072-1092 ^d ID NO. 1999

a Sequences from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

^c These sequences were aligned to derive the corresponding primer.

d The nucleotide positions refer to the C. albicans tuf sequence fragment (SEQ ID NO. 2004).

Annex II: Specific and ubiquitous primers for nucleic acid amplification (atpD sequ nc s).

				Originating	DNA fragment
5	SEQ ID NO.	Nucleotide	sequence	SEQ ID NO.	Nucleotide position
10	Bacterial	species:	Acinetobacter baumanni	i	
	1690	5'-CAG GTC	CTG TTG CGA CTG AAG AA	243	186-208
	1691 ^b	5'-CAC AGA	TAA ACC TGA GTG TGC TTT C	243	394-418
15	Bacterial	species:	Bacteroides fragilis		
	2134	5'-CGC GTG	AAG CTT CTG TG	929	184-200
	2135 ^b	5'-TCT CGC	CGT TAT TCA GTT TC	929	395-414
20	Bacterial	species:	Bordetella pertussis		
	2180	5'-TTC GCC	GGC GTG GGC	1672 ^C	544-558
	2181 ^b	5'-AGC GCC	ACG CGC AGG	1672 ^C	666-680
25	Bacterial	species:	Enterococcus faecium		
	1698	5'-GGA ATC	AAC AGA TGG TTT ACA AA	292	131-153
	1699 ^b	5'-GCA TCT	TCT GGG AAA GGT GT	292	258-277
30	1700	5'-AAG ATG	CGG AAA GAA GCG AA	292	271-290
	1701 ^b	5'-ATT ATG	GAT CAG TTC TTG GAT CA	292	439-461
	Bacterial	species:	Klebsiella pneumoniae		
35	1331	5'-GCC CTT	GAG GTA CAG AAT GGT AAT GAA	GTT 317	88-118
	1332 ^b	5'-GAC CGC	GGC GCA GAC CAT CA	317	183-203

a These sequences were aligned to derive the corresponding primer.

⁴⁰ b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

C Sequence from databases.

PCT/CA00/01150 WO 01/23604

Specific and ubiquitous prim rs for nucleic acid Annex II: amplification (atpD sequences).

-										Originating I	NA fragment
:	SEQ ID NO.	Nucleotide	sequ	ence	:					SEQ ID NO.	Nucleotide position
-	Bacterial spe	cies:	St.	rept	oco	ccu	s ag	rala	cti	ae	
	627	5'-ATT GTC	TAT	AAA	AAT	GGC	GAT	AAG	тC	379-383 ^a	42-67 ^b
	625 ^C	5'-CGT TGA								379-383 ^a	206-231 ^b
		5'-AAA ATG	ccc	እጥአ	እርጥ	C Δ C	ΔΔΔ	AAG	TΑ	379-383 ^a	52-77 ^b
	628 625 ^C	5'-AAA ATG								379-383 ^a	206-231 ^b
										379-383 ^a	42-67 ^b
	627 626 ^C	5'-ATT GTO 5'-TAC CAO	COT	AAA TTT A	AAT	AAG	GTG	CTA	AT	379-383 ^a	371-396 ^b
	6260										52-77 ^b
	628	5'-AAA ATG								379-383 ^a 379-383 ^a	-
	626 ^C	5'-TAC CA									3,1 330
	Bacterial gro	oup:	Ca	mpy.	loba	icte	r j	eju	i i	and C. coli	
	2131	5'-AAG CM	A TTG	TTG	TAA	TTA	TTG	AAA	G	1576,1600, 1849,1863,2139	7-31 ^e 9d,a
	2132 ^c	5'-TCA TA	T CCA	TAG	CAA	TAG	·TTC	TA		1576,1600, 1849,1863,213	92-114 ^e
	Bacterial qe	nus:	В	orđe	tel	la s	p.				
		5'-ATG AG	C 3DC	י ככז	ACC	י אייר	י כיתיו	י ראַפּ	TG:	1672 ^d	1-26
	825 826 ^C	5'-ATG AG									1342-1367
	. <u>Fungal genus</u>	:	Ci	anaı	da :	вр.					_
	634	5'-AAC AC	Y GTO	C AGF	RCI	TA	GC?	ATC	G GA	460-472, 474-478 ^a	101-126 ^f
	635 ^C	5'-AAA CC	R GT	I ARI	R GCF	R ACT	r CTI	GC1	CI		617-642 ^f

a These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the S. agalactiae atpD sequence fragment 45 (SEQ ID NO. 380).

C These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

d Sequence from databases.

 $^{^{\}mathrm{e}}$ The nucleotide positions refer to the C. jejuni atpD sequence fragment (SEQ ID NO. 1576).

f The nucleotide positions refer to the C. albicans atpD sequence fragment (SEQ ID NO. 460).

PCT/CA00/01150 WO 01/23604

Specific and ubiquitous primers for nucleic acid Annex II: amplification (atpD sequences) (continued).

5			Originating D	NA fragment
	SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
0		Universal primers		-
5	562	5'-CAR ATG RAY GAR CCI CCI GGI GYI MGI ATG	243,244,262, 264,280,284, 291,297,309, 311,315,317, 324,329,332, 334-336,339, 342,343,351, 356,357,364- 366,370,375,	
25	563 ^C	5'-GGY TGR TAI CCI ACI GCI GAI GGC AT	379,393 ^a 243,244,262, 264,280,284, 291,297,309, 311,315,317, 324,329,332, 334-336,339, 342,343,351, 356,357,364	
30			366,370,375 379,393 ^a	
35	564	5'-TAY GGI CAR ATG AAY GAR CCI CCI GGI AA	243,244,262 264,280,284 291,297,309 311,315,317 324,329,332 334-336,339	, , ,
40			342,343,351 356,357,364 366,370,375 379,393 ^a	<u>,</u>
45 50	565 ^c	5'-GGY TGR TAI CCI ACI GCI GAI GGD AT	243,244,262 264,280,284 291,297,309 311,315,317 324,329,332 334-336,339 342,343,351 356,357,364	, , ,
			366,370,375 379,393 ^a	

a These sequences were aligned to derive the corresponding primer. , 55

b The nucleotide positions refer to the K. pneumoniae atpD sequence fragment (SEQ ID NO. 317).

^C These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex II: Specific and ubiquitous primers for nucleic acid amplification (atpD sequences) (continued).

			Originating DNA fragment
	SEQ ID NO.	Nucleotide sequence	SEQ ID Nucleotide NO. position
		Universal primers (continued)	
	640	5'-TCC ATG GTI TWY GGI CAR ATG AA	248,284,315, 513-535 ^b 317,343,357,
			366,370,379,393 ^a
	641 ^C	5'-TGA TAA CCW ACI GCI GAI GGC ATA C	317,343,357,
			366,370,379,393 ^a
	642	5'-GGC GTI GGI GAR CGI ACI CGT GA	248,284,315, 438-460 ^b 317,343,357,
			366,370,379,393 ^a
	641 ^C	5'-TGA TAA CCW ACI GCI GAI GGC ATA C	G 248,284,315, 684-709 ^b 317,343,357,
			366,370,379,393 ^a
		Sequencing primers	
	566	5'-TTY GGI GGI GCI GGI GTI GGI AAR A	AC 669 ^d 445-470
	567 ^C	5'-TCR TCI GCI GGI ACR TAI AYI GCY T	
)			ac 669 ^d 445-470
	566	5'-TTY GGI GGI GCI GGI GGI AAR A	666d 901-920
	814	5'-GCI GGC ACG TAC ACI GCC TG	666-
	568	5'-RTI ATI GGI GCI GTI RTI GAY GT	669 ^d 25-47
5	567 ^C	5'-TCR TCI GCI GGI ACR TAI AYI GCY	rg 669 ^d 883-908
		a cor cor cor por chy cm	672 ^d 31-53
	570	5'-RTI RYI GGI CCI GTI RTI GAY GT 5'-TCR TCI GCI GGI ACR TAI AYI GCY	
	567 ^C	5'-TCR TCT GCT GGT ACR TAI ATT GCT	
)	572	5'-RTI RTI GGI SCI GTI RTI GA	669 ^d 25-44
	567 ^C	5'-TCR TCI GCI GGI ACR TAI AYI GCY '	TG 669 ^d 883-908
	5.66	5'-RTI RTI GGI SCI GTI RTI GAT AT	671 ^d 31-53
	569 567 ^C	5'-TCR TCI GCI GGI ACR TAI AYI GCY	
5	201-		
	571	5'-RTI RTI GGI CCI GTI RTI GAT GT	670 ^d 31-53
	567 ^C	5'-TCR TCI GCI GGI ACR TAI AYI GCY	TG 669 ^d 883-908

on a These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the *K. pneumoniae atpD* sequence fragment (SEQ ID NO. 317).

These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

⁵⁵ d Sequences from databases.

Annex II: Specific and ubiquitous primers for nucleic acid amplification (atpD sequences) (continued).

			Originating DNA fragment
5	SEQ ID NO.	Nucleotide sequence	SEQ ID Nucleotide NO. position
10		Sequencing primers (continued)	
	700	5'-TIR TIG AYG TCG ART TCC CTC ARG	669 ^a 38-61
	567 ^b	5'-TCR TCI GCI GGI ACR TAI AYI GCY TG	669 ^a 883-908
15	568	5'-RTI ATI GGI GCI GTI RTI GAY GT	669 ^a 25-47
13	573 ^b	5'-CCI CCI ACC ATR TAR AAI GC	666 ^a 1465-1484
	574	5'-ATI GCI ATG GAY GGI ACI GAR GG	666 ^a 283-305
	573 ^b	5'-CCI CCI ACC ATR TAR AAI GC	666 ^a 1465-1484
20 .	574	5'-ATI GCI ATG GAY GGI ACI GAR GG	666 ^a 283-305
	708 ^b	5'-TCR TCC ATI CCI ARI ATI GCI ATI AT	666 ^a 1258-1283
	681	5'-GGI SSI TTY GGI ISI GGI AAR AC	685 694-716
25	682 ^b	5'-GTI ACI GGY TCY TCR AAR TTI CCI CC	686 1177-1202
	681	5'-GGI SSI TTY GGI ISI GGI AAR AC	685 694-716
	683p	5'-GTI ACI GGI TCI SWI AWR TCI CCI CC	685 1180-1205
30	681 699	5'-GGI SSI TTY GGI ISI GGI AAR AC 5'-GTI ACI GGY TCY TYR ARR TTI CCI CC	685 694-716 686 1177-1202
	681	5'-GGI SSI TTY GGI ISI GGI AAR AC	685 694-716
35	812 ^b	5'-GTI ACI GGI TCY TYR ARR TTI CCI CC	685 1180-1205
33	1213	5'-AAR GGI GGI ACI GCI GCI ATH CCI GG	714 ^a 697-722
	1212 ^b	5'-CCI CCI RGI GGI GAI ACI GCW CC	714 ^a 1189-1211
	1203	5'-GGI GAR MGI GGI AAY GAR ATG	709 ^a 724-744
40	1207 ^b	5'-CCI TCI TCW CCI GGC ATY TC	709 ^a 985-1004
	1204	5'-GCI AAY AAC ITC IWM YAT GCC	709 ^a 822-842
	1206 ^b	5'-CKI SRI GTI GAR TCI GCC A	709 ^a 926-944
45	1205	5'-AAY ACI TCI AWY ATG CCI GT	709 ^a 826-845
	1207 ^b	5'-CCI TCI TCW CCI GGC ATY TC	709 ^a 985-1004
	2282	5'-AGR RGC IMA RAT GTA TGA	714 ^a 84-101
50	2284b	5'-TCT GWG TRA CIG GYT CKG AGA	714 ^a 1217-1237
50	2283	5'-ATI TAT GAY GGK ITT CAG AGG C	714 ^a 271-292
	2285 ^b	5'-CMC CIC CWG GTG GWG AWA C	714 ^a 1195-1213

^{. 55} a Sequences from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex III: Internal hybridization probes for specific detection of tuf sequences.

				Originating DN	A fragment
5	SEQ ID NO.	Nucleotide	sequence		Nucleotide position
10	Bacterial sp	ecies:	Abiotrophia adiacen	s	
•	2170	5'-ACG TGA	CGT TGA CAA ACC A	1715	313-331
15	Bacterial sp	ecies:	Chlamydia pneumonia	e	
13	2089 2090		AAC TTA TTG ACC TT TGG AGT CGA AAT G	20 20	136-155 467-485
:	Bacterial sp	ecies:	Enterococcus faecal	is	
	580 603 1174	5'-GGT ATT	CCA GCT ACA ATC ACT CCA AAA GAC GAA ACA TC GGT GAA GTT CGC	C 62-63,607 ^a 62-63,607 ^a 62-63,607 ^a	440-459 ^b
25	Bacterial sr	oecies:	Enterococcus faeciu	un.	
	602	5'-AAG TTC	G AAG TTG TTG GTA TT	64,608 ^a	426-445 ^C
:	Bacterial sp	pecies:	Enterococcus gallir	narum	
30	604	5'-GGT GAT	GAA GTA GAA ATC GT	66,609 ^a	419-438 ^d
,	Bacterial s	pecies:	Escherichia coli		
35	579	5'-GAA GG	C CGT GCT GGT GAG AA	78	503-522
	2168	5'-CAT CA	A AGT TGG TGA AGA AGT TG	78	409-431
40	Bacterial s	oecies:	Neisseria gonorrhoe	eae	
40	2166	5'-GAC AA	A CCA TTC CTG CTG	126	322-339 ^e
	Fungal spec	ies:	Candida albicans		
، 45	577	5'-CAT GA	T TGA ACC ATC CAC CA	.407-411 ^a	406-425 ^f
	Fungal spec	<u>ies</u> :	Candida dubliniens:	is	
: 50	578	5'-CAT GA	T TGA AGC TTC CAC CA	412,414-415 ^a	418-437 ^g
	a These sequen b The nucleoti 607).	ces were align de positions	ed to derive the correspond refer to the E. faecalis t	ing primer. uf sequence fragment	(SEQ ID NO.
: 55	C The nucleoti	de positions	refer to the E. faecium to	of sequence fragment	(SEQ ID NO.
	d The nucleoti		refer to the E. gallinarum		
1	e The nucleoti		refer to the N. gonorrhoeae		
50	f The nucleoti		refer to the C. albicans tu	·	
	g The nucleoti ID NO. 414).		refer to the C. dubliniensis	s tuf(EF-1) sequence	rragment (SEQ

Annex III: Internal hybridization probes for specific detection of tuf sequences (continued).

			C	riginating D	NA fragmen
SEQ ID NO.	Nucleotide sequence		_	SEQ ID	Nucleotide position
Bacterial s	pecies: Haemoph	ilus inf	luenzae		
581	5'-ACA TCG GTG CAT TA	AT TAC GTG	G	610 ^a	551-572
Bacterial s	pecies: Mycopla	sma pneu	moniae		
2095	5'-CGG TCG GGT TGA AG	CG TGG		2097 ^a	687-704
Bacterial s	species: Staphyl	ococcus	aureus		
584	5'-ACA TGA CAC ATC TA	AA AAC AA		176-180 ^b	369-388 ^C
585	5'-ACC ACA TAC TGA A'			176-180 ^b	525-544 ⁰
586	5'-CAG AAG TAT ACG TA			176-180 ^b	545-564 ⁰
587	5'-CGT ATT ATC AAA A			176-180 ^b	555-5749
588	5'-TCT TCT CAA ACT A			176-180 ^b	593-6129
Bacterial	species: Staphyl	ococcus	epidermi	dis	
		** **C **		185,611 ^b	445-464 ⁰
589	5'-GCA CGA AAC TTC T 5'-TAT ACG TAT TAT C			185,611 ^b	627-646°
590	5'-TCC TGG TTC TAT T			185,611 ^b	586-605
591	5'-CAA AGC TGA AGT A			185,611 ^b	616-635
592 593	5'-TTC ACT AAC TAT C			185,611 ^b	671-690
Bacterial			haemolyt	icus	
				186,188-190 ^b	2 437-456
594	5'-ATT GGT ATC CAT G			186,188-190 ^k	
595	5'-TTA AAG CAG ACG T			100,100-190	013 034
<u>Bacterial</u>	species: Staphy	lococcus	hominis		
596	5'-GAA ATT ATT GGT A	TC AAA GA		191,193-196 ¹	431-450
597	5'-ATT GGT ATC AAA			191,193-196 ¹	437-456
598	5'-AAT TAC ACC TCA			191,193-196 ¹	595-614

a Sequences from databases.

45

b These sequences were aligned to derive the corresponding probe.

 $^{^{\}rm C}$ The nucleotide positions refer to the S. aureus tuf sequence fragment (SEQ ID NO. 179).

^{, 50} d The nucleotide positions refer to the S. epidermidis tuf sequence fragment (SEQ ID NO. 611).

e The nucleotide positions refer to the S. haemolyticus tuf sequence fragment (SEQ ID NO. 186).

f The nucleotide positions refer to the *S. hominis tuf* sequence fragment (SEQ ID NO. 191).

Annex III: Internal hybridization probes for specific detection of tuf sequences (continued).

-									_		0	riginating 1	DNA fragment
5	SEQ ID N	ю.	Nucleot	ide	sequ	ence	:					SEQ ID NO.	Nucleotide position
-	Bacter:	ial sp	ecies:		Sta	aphy	100	occ	us s	sapr	ophyt	icus	
	599		5 ′ -CGG	тса	AGA	ААТ	CGA	AAT	CA			198-200 ^a	406-425 ^b
	600		5'-ATG									198-200 ^a	431-450 ^b
	601		5'-GTT									198-200 ^a	536-555 ^b
	695		5'-GTT									198-200 ^a	563-582 ^b
	Bacter	ial sp	ecies:		St	rep	tocc	occu	s ag	gala	ctiae	9	
	- o - C	5	CAA CTT	CCT	CCT	ጥር እ	CAC	CAA	CAG	т		207-210 ^a	404-431 ^d
	5820	5'-TT	CAA CTT	CGI	CGI	עטעע	CAL	CUM	TAA	TAC	CAA C	G 207-210 ^a	433-467 ^d
	583° 1199	5'-CAA 5'-GTA	TTA AAG	AAG	ATA	TCC	AAA	AAG	C	0	J V	207-210 ^a	438-462 ^d
			ecies:							new	nonia	e	
	1201		5′-TCA	AAG	AAG	AAA	CTA	. AAA	дAG	CTG	T	971,977, 979,986 ^a	513-537 ^e
	Bacter	ial sr	ecies:		St	rep	toc	occi	s p	yoge	enes		
	1200		5'-TCA	AAG	AAG	AAA	CTA	AAA	AAG	CTG	T	1002	473-497
	Bacter	ial or	our.		En	ter	000	ccus	ca	sse.	lifla	vus-flave	scens-
	Bacter	ıaı gı	<u>.oup</u> .						rou				
	500		5′-ATT		ר ככא	ጥጥር	י כיתי	\ CG1				58.65.66ª	527-544 [£]
	620 1122		5'-TG									58,65,66ª	
	Bacter	ial g								eme.	lla s	p., A. ad	iacens
	2172		5′-GT(G TT	g aaa	TG	r TC	C GT	AA A			58-62,67-7 87-88,607-6 727,871	L, 477-496 ^g 09,
												1715,1722	a

⁴⁵ a These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the S. saprophyticus tuf sequence fragment (SEQ ID NO. 198).

C These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

 $^{^{}m d}$ The nucleotide positions refer to the S. agalactiae tuf sequence fragment (SEQ ID NO. 209).

e The nucleotide positions refer to the *S. pneumoniae tuf* sequence fragment (SEQ ID NO. 986).

f The nucleotide positions refer to the E. flavescens tuf sequence fragment (SEQ ID NO. 65).

⁹ The nucleotide positions refer to the $\it E.~faecium~tuf~sequence~fragment~(SEQ~ID~NO.~608)$.

Annex III: Internal hybridization probes for specific detection of tuf sequences (continu d).

				Originating D	NA fragment
SEQ ID NO.	Nucleotide	sequence		SEQ ID NO.	Nucleotide position
Bacterial ge	enus:	Gemella			
2171	5'-TCG TTG	GAT TAA CTG	AAG AA	87,88 ^a	430-449b
Bacterial ge	enus:	Staphyloc	coccus sp.		
605	5'-GAA ATG	TTC CGT AAA	TTA TT	176-203 ^a	403-422 ^C
606		CTA CGC TGA		176-203 ^a	420-439 ^C
1175		GGT GTA GAA		176-203 ^a	391-411 ^C
1176		TGT AGA AAT		176-203 ^a	393-411 ^C
Bacterial ge	enus:	Streptoco	occus sp.		
1202	5'-GTG TTG	AAA TGT TCC	GTA AAC A	206-231,971, 977,979,982 - 98	
Fungal spec	ies:	Candida a	albicans		
1156	5'-GTT GAA	ATG CAT CAC	GAA CAA TT	407-412,624 ⁶	680-702 ^e
Fungal grou	ο:	Candida a	albicans and	l C. tropicali	s
1160	5'-CGT TTC	: TGT TAA AGA	AAT TAG AAG	407-412, 429,624 ^a	748-771 ^e
Fungal spec	ies:	Candida	dubliniensi	5	
1166	ፍ / - አርር ጥ ጥ	A AGA ATG TTI	r crg rca a	414-415 ^a	750-771 [£]
1168		TTG GTT GA		414-415 ^a	_
Fungal spec	<u>ies</u> :	Candida	glabrata		
1158 1159	5'-AAG AGG 5'-TGA AGG	TAA TGT CTC	G TGG T G TGA	417 417	781-799 718-735
Fungal spec	ies:	Candida	krusei		
	5'-TCC AG			422	720-737

⁵⁰ a These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the *G. haemolysans tuf* sequence fragment (SEQ ID NO. 87).

c The nucleotide positions refer to the S. aureus tuf sequence fragment (SEQ ID NO. 179).

d The nucleotide positions refer to the S. pneumoniae tuf sequence fragment (SEQ ID NO. 986).

e The nucleotide positions refer to the C. albicans tuf(EF-1) sequence fragment (SEQ ID NO. 408).

f The nucleotide positions refer to the C. dubliniensis tuf(EF-1) sequence fragment (SEQ ID NO. 414).

Annex III: Internal hybridization prob s for specific detection of tuf sequences (continued).

					Originating DN	A fragment
SI	EQ ID NO.	Nucleotide	sequence		SEQ ID NO.	Nucleotide position
F	ungal group:		Candida lus:	itaniae and	C. guillermo	ondii
	1162	5'-CAA GTC	CGT GGA AAT GCA	4	418,424 ^a	682-699 ^b
F	ungal specie	<u>s</u> :	Candida para	apsilosis		
	1157	5'-AAG AAC	GTT TCA GTT AAG	G GAA AT	426	749-771
F	<u>'ungal specie</u>	<u>s</u> :	Candida zey	lanoides		
	1165	5'-GGT TTC	AAC GTG AAG AAG	C	432	713-730
<u>F</u>	<u>Sungal genus:</u>		Candida sp.			
	1163	5'-GTT GGT	TTC AAC GTT AA	G AAC	407-412,414- 415,417,418, 422,429 ^a	728-748 ^C
	1164	5'-GGT TTC	AAC GTC AAG AA	С	413,416,420, 421,424,425, 426,428,431 ^a	740-757 ^b
	1167	5'-GTT GGT	TTC AAC GT		406-426, 428- 432, 624 ^a	728-741 ^C

a These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the C. lusitaniae tuf(EF-1) sequence fragment (SEQ ID NO. 424).

 $^{^{\}rm C}$ The nucleotide positions refer to the $^{\rm C.}$ albicans tuf(EF-1) sequence fragment (SEQ ID NO. 408).

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Accession

SEO ID

(F-Strategy for the selection of amplification/sequencing primers from atpD type) sequences. Annex IV:

	2	O E C C C C C C C C C C C C C C C C C C	91 GTGT ACGTCCCTGC GGACGACT	910 NO.:	#: X7687X
	2	hgrgcal cgccccctr arcsacctsGTGTTCG issis Acrgcal cgcccccrc GrgcalarTCTGTTCG inosa Aaarcal cgcccccrc arcsacctcGTGTTCG inosa Aaarcal cgcccccrc arcsacctCGTGTTCG	COTGT ACGTGCCTGC CGACGACT COTAT ACGTTCCCGC GGACGACC AGTAT ACGTACCTGC GGATGACT	1 1 1 1	Genome project Genome project J01594 Genome project
	2		NATOT ATGTGCCGGC CGACGACC TATOT ACGTGCCCGC CGACGACC GGTCT ACGTGCCCGC CGACGACC	1 1 1	U64318 X76879 Z73419
	15	GGGTCAC TGGGCCCGTC GTCGACGTCGTGTTCG AGGDAAT TGGCCCTGTG GTCGATGTGTTGTTTG AAATTAT TGGCCCAGIT ATAGATGTGGTATTTG	IGITI ACGIACCGGC IGAIGACT 3GITI ACGIACCIGC GGAIGAIT 3GITI ACGAICCAGC CGAIGACT	672	M2224/ M22535 U10505
		icum	IGTAT ATGITCCTGC TGATGACC GATCT ATGIGCCAGC TGATGACT	671	AF101055 U43738 AF004014
	20	M. pylori H. pylori	19199		
2		Selected sequences for universal primers		568	
271	25	25		5 / 2 5 6 9 5 7 1	
•		RIERT IGGICCIGTI REIGATGT TING GIGGICCIGG IGTIGGIAAR AC		999	
	30	Selected sequence for universal primer*	CA RGCIRTIT AYGTICCIGC IGAYGA	567	
	35	The sequence numbering refers to the <i>Escherichia coli atpD</i> gene fragment (SEQ ID NO. selected sequences or match those sequences. Mismatches are indicated by lower-case letter	Nucleotides in capitals s indicate gaps in the se	are id quences	intical to the displayed.
	1			, D	"M" stands for

This sequence is the reverse-complement of the selected primer.

9

to any of the four nucleotides A, C, G or T.

"R" "Y" "M" "K" "W" and "S" designate nucleotide positions which are degenerated. "R" stands for A or G; "Y" stands for C or T; "M" stands for C or G. "I" stands for inosine which is a nucleotide analog that can bind A or C; "K" stands for G or T; "W" stands for A or T; "S" stands for C or G. "I" stands for Inosine which is a nucleotide analog that can bind

niversal amplification/sequencing primer:		
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	serectron	sednences.
	the	ype)
(for th	(V-t
	Strategy	from atpD (V-type)
	Annex V:	

SEQ ID NO.: 685 687 693 688 692 689 689	681	682 683
CC AGGRCCETT GGTGCAGGGA AGACAGTTCTGGTGGAG ATATCTCTGA ACCAGTGACT CA CC AGGRCCGTT GGGTCGGGA AGACAGTTCTGGTGGAG ATATCTCTGGA ACCAGTGACT CA CC GGGGCCGTTC GGGTCGGGA AGACGGTCCGGGCGGCG ACTTCTCGGA GCCGGTGACC CA CC TGGGGCCTTC GGATGTGGCA AGACGTCCGGGGGGAG ACTTCTCAGAG CCCGTGACG AC CC TGGGGCCTTT GGATGCGGAA AGACGGTCCTGGAGGTG ACTTTCTCAGA CCCTGAGGG C CC TGGCGCGTTT GGTGTGGAA AAACTTGCCTGGAGGTG ACTTTCTCAGA CCCTGTAACT AC CC TGGTGCATTT GGTGTGGGAA AAACAGTGCAGGGGGAA ACTTTGAAGA ACCAGTGACT CA	GGISSITTY GGIISIGGIA ARAC	GGIGGIA AYTTYGARGA RCCIGTIAC GGIGGIG AYWTIWSIGA ICCIGTIAC
E. hirae H. salinarum T. thermophilus Human T. congolense P. falciparum C. pneumoniae	15 Selected sequences for universal primers	Selected sequences
5 10	15	6

"R" "Y" "M" "K" "W" and "S" designate nucleotide positions which are degenerated. "R" stands for A or G; "Y" stands for C or T; "W" stands for B or T; "W" stands for B or T; "W" stands for B or T; "W" stands for G or T; "W" stands for B or T; "W" stand case letters. Mismatches for SEQ ID NO. 683 are indicated by underlined nucleotides. Dots indicate gaps in the sequences identical to the selected sequences or match those sequences. Mismatches for SEQ ID NOs. 681 and 682 are indicated by lowerdisplayed. 25

The sequence numbering refers to the Enterococcus hirae atpD gene fragment (SEQ ID NO. 685). Nucleotides in capitals are

a nucleotide analog that can bind to any of the four nucleotides A, C, G or T. 30

These sequences are the reverse-complement of the selected primers.

for universal primers^a

of universal amplification/sequencing (organelle origin). sednences selection primers from tuf (M) the for Strategy Annex VI:

5 C. necformanse Andra Caractece Aspertance Charactece Aspertance Conception Andra Caractece Conception Andra Caractece Conception Andra Caractece Conception Conception Andra Caractece Conception Conception Caractece Caractece Conception Caractece Caractece Conception Caractece Caracte	sion	03 79 33 58 58 00 125 521 28 27			des in nd 664 . Dots	G; "Y" G. "I"	
5. c. neoformans* AAGAA CATGATTACT GATGATCTC. AGGATGAGA GAGATGGAGA GAGATGAGA GAACGATTGC GT - 665 S. c. neoformans* AAGAA CATGATTACT GATGATCTC. AGGATGACTC. GCCTGTCAGA GAGATGAGC TTG - 665 S. c. neoformans* AAGAA CATGATTACT GATGATCTC. AGGATGACTC. GCCTGTCAGA GAGATGAGC TTG - 665 S. c. neoformans* AAGAA CATGATTACT GATGATCTC. AGGATGAGGG GALGATTGG GT - 670 N. volvulus* AAGAA CATGATTACA GGGGATCTC AGGATGAGGG GALGATTGG GT - 670 G. max B2P AAAA ATGATTACA GGGGACCCC AAATGAGGG. "GCCTATTAGA GAGAGGGAGA AAACGATTGG GT - 670 E. c. neoformans* AAGAA CATGATCAC GGGGCGCCC AAATGAGGGG. "GCCATTAGA GAGAGGGAGA AAACGATTGG GT - 670 E. c. neoformans* AAGAA CATGATCAC GGGGCGCCC AAATGAGGGG. "GCCATTAGA GAGAGGAGA AAACGATTGG GT - 670 E. c. neoformans* AAGAA TATGATTACA GGGGCGCCC AAATGAGTGG. "GCCATTAGA GAGAGGAGA AAACGATTGG GT - 670 E. c. neoformans* AAGAA TATGATTACA GGGGCGCCC AAATGAGTGG. "GCCATTAGA GAGAGGAGA AAACGATTGG GT - 670 E. c. neoformans* AAGAA TATGATTACA GAGACGACCACA AAATGAGTGG. "GCCATTAGA GAGAGGAGA AAACGATTGG GT - 670 E. c. neoformans* AAAA TATGATTACA GAGACGACCACA AAATGAGTGG. "GCCATTAGA GAGAGGAGA AAACGATTGG GT - 670 E. c. neoformans* AAAA TATGATTACA GAGACGCCCC AAATGAGTGG. "GCCATTAGA GAGAGGAGA AAACGATTGG GT - 670 E. c. neoformans* AAAA TATGATTACA GAGACGCCC AAATGAGTG. "GCCATTAGA GAGAGGAGA AAACGATGG GT - 670 E. c. neoformans* AAAA TATGATTACA GAGACGCCC AAATGAGTG. "GCCATTAGA GAGAGGAGA AAACGATGG GT - 670 E. c. neoformans* AAAA TATGATTACA GAGACGCCC AAATGAGTG. "GCCATTAGA GAGAGGAGA AAACGATGG GT - 670 E. c. neoformans* AAAA TATGATTACA GAGACGCCC AAATGAGTG. "GCCATTAGAGGGGAA AAACGATGG GT - 670 E. c. neoformans* AAAAA TATGATTACA GAGAGGACA AAATGAGTG "GCCATTAGAGGGGAA AAAGAGATG - 670 E. c. neoformans* AAAAA TATGATTACA GAGAGGAGA AAAAGATG - 670 E. c. neoformans* AAAAAA TATGATTACA GAGAGGAGAA AAAAGATG - 670 E. c. neoformans* AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	Acces.	U81803 X00779 M64333 X03558 Y15107 Y15108 - AF007125 AI755521 Y11431 K00428			leotic 652 a otides		
5 C. neoformans ^a AAGAA CATGATCACC GCTaCCCCC AGGCTGACGA GAGGAGA (A. o. volvalus ^a AAGAA CATGATCACC GCTACCTCC AGGCTGACTGA GAGGA (A. o. volvalus ^a AAGAA CATGATCACA GGTACTCCC AGGCTGACTGA. TGCTGTGGCT GAGGA (A. o. volvalus ^a AAGAA CATGATCACA GGGCACTCC AGGCTGACTGA. TGCTGTTGGCT GAGGA (A. o. volvalus ^a AAGAA CATGATCACA GGGCACTCC AGGCTGACTGA. TGCTGTTGGCT GAGGA (A. o. volvalus ^a AAGAA CATGATCACC GGGCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA CATGATCACC GGCCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA CATGATCACC GGCCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA CATGATCACC GGTGCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTAGAG GAAGA AAAAA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTAGAG GAAGA TATGATTACA TGCTATTAGAGA TATGATTACA TGCTATTAGAGA TATGATTACA TATGATTAC	SEQ ID	665	664	652 561	. Nuc NOs. nuclec	for is for	
5 C. neoformans ^a AAGAA CATGATCACC GCTaCCCCC AGGCTGACGA GAGGAGA (A. o. volvalus ^a AAGAA CATGATCACC GCTACCTCC AGGCTGACTGA GAGGA (A. o. volvalus ^a AAGAA CATGATCACA GGTACTCCC AGGCTGACTGA. TGCTGTGGCT GAGGA (A. o. volvalus ^a AAGAA CATGATCACA GGGCACTCC AGGCTGACTGA. TGCTGTTGGCT GAGGA (A. o. volvalus ^a AAGAA CATGATCACA GGGCACTCC AGGCTGACTGA. TGCTGTTGGCT GAGGA (A. o. volvalus ^a AAGAA CATGATCACC GGGCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA CATGATCACC GGCCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA CATGATCACC GGCCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA CATGATCACC GGTGCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTAGAG GAAGA AAAAA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTAGAG GAAGA TATGATTACA TGCTATTAGAGA TATGATTACA TGCTATTAGAGA TATGATTACA TATGATTAC		CGT TGT AGC CGC CGC CGC CGC CGC CGC CGC CGC CGC		υ ,	619) SEQ ID Lined	stands stand s A, C	
5 C. neoformans ^a AAGAA CATGATCACC GCTaCCCCC AGGCTGACGA GAGGAGA (A. o. volvalus ^a AAGAA CATGATCACC GCTACCTCC AGGCTGACTGA GAGGA (A. o. volvalus ^a AAGAA CATGATCACA GGTACTCCC AGGCTGACTGA. TGCTGTGGCT GAGGA (A. o. volvalus ^a AAGAA CATGATCACA GGGCACTCC AGGCTGACTGA. TGCTGTTGGCT GAGGA (A. o. volvalus ^a AAGAA CATGATCACA GGGCACTCC AGGCTGACTGA. TGCTGTTGGCT GAGGA (A. o. volvalus ^a AAGAA CATGATCACC GGGCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA CATGATCACC GGCCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA CATGATCACC GGCCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA CATGATCACC GGTGCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTAGAG GAAGA AAAAA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTAGAG GAAGA TATGATTACA TGCTATTAGAGA TATGATTACA TGCTATTAGAGA TATGATTACA TATGATTAC		cGTTGc aGTTGc aGTTGc TGTTGG cGTTGG cGTTGG cGTTGG cGTTGG		TRIWG IGI ^d	ID NO.	"R" s T; "S" eotide	
5 C. neoformans ^a AAGAA CATGATCACC GCTaCCCCC AGGCTGACGA GAGGAGA (A. o. volvalus ^a AAGAA CATGATCACC GCTACCTCC AGGCTGACTGA GAGGA (A. o. volvalus ^a AAGAA CATGATCACA GGTACTCCC AGGCTGACTGA. TGCTGTGGCT GAGGA (A. o. volvalus ^a AAGAA CATGATCACA GGGCACTCC AGGCTGACTGA. TGCTGTTGGCT GAGGA (A. o. volvalus ^a AAGAA CATGATCACA GGGCACTCC AGGCTGACTGA. TGCTGTTGGCT GAGGA (A. o. volvalus ^a AAGAA CATGATCACC GGGCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA CATGATCACC GGCCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA CATGATCACC GGCCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA CATGATCACC GGTGCCCCCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCC AGATGGACGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCCGCCA AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTGAGA GAAGA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTAGAG GAAGA AAAAA TATGATTACA GGACCTGCCC AAATGGATGG. TGCTATTAGAG GAAGA TATGATTACA TGCTATTAGAGA TATGATTACA TGCTATTAGAGA TATGATTACA TATGATTAC					(SEQ atchesed by	ated. A or	
C. neoformans ^a AAGAA CATGATCACC GGTACCTCCC AGGCTGACTGCGCCgr. S. cerevisiae ^a AAGAA CATGATTACT GGTACTCTCT AAGCTGACTGCGCTgr. O. volvulus ^a AAGAA CATGATTACT GGTACTCTCTC AGGCTGACTGTGCTgr. AAGAA CATGATTACA GGGACACTC AGGCTGACTGTGCTgr. AAGAA CATGATTACA GGGACACCCCC AGATGCACGGTGCTgr. AAGAA CATGATCAC GGCGCCCCCC AGATGCACGGCGCAS. AAGAA CATGATCAC GGCGCCCCCC AGATGCACGGCGCAS. AAGAA CATGATCAC GGCGCCCCCC AGATGCACGGCGCAS. AAGAA CATGATTACA GGACCACCCC AGATGCACGGCGCAS. AAGAA CATGATTACA GGACCACCCC AGATGCACGGCGCAS. AAGAA TATGATTACA GGACCACCAC AAATGCACGGCGCAS. AAGAA TATGATTACA GGACCACCAC AAATGCACGGCGCAS. AAGAA TATGATTACT GGACCTCCCC AAATGCACGGTGCTAS. A thalianab AAGAA TATGATTACT GGACCTCCCC AAATGCACGGTGCTAS. A thalianab AAAA TATGATTACT GAACTCCCCC AAATGCACGGTGCTAS. A thalianab AAAAA TATGATTACT GAACTCCCCC AAATGCACGGTGCTAS. A thalianab AAAAA TATGATTACT GAACTCCCCC AAATGCACGGTGCTAS. A thalianab AAAAA TATGATTACT AAATGCATCCCC AAATGCACGGTGCTAS. A thalianab AAAAA TATGATTACT AAATGCATCCCCC					(M) gene nces. Mism re indicat	re degener tands for f the four	
C. neoformansa AAGAA CATGATTACT GGTACCTCCC AGGCTGATG S. cerevisiaea AAGAA TATGATTACA GGTACTTCTC AAGCTACTG O. volvulus AAGAA TATGATTACA GGTACTTCTC AGGCTGATG Humana AAGAA TATGATTACA GGTACTTCTC AGGCTGATG 10 G. max Blb AAGAA CATGATTACA GGGCACTCC AGATGGANGG E. coli G. max Blb AAGAA CATGATTACA GGGCCTGCCC AGATGGANGG E. coli G. max Blb AAGAA CATGATTACA GGGCCTGCCC AGATGGANGG E. coli G. max Blb AAGAA CATGATTACA GGGCCTGCCC AGATGGANGG E. coli G. max Blb AAGAA CATGATTACA GGGCCTGCCC AGATGGANGG E. coli G. max Blb AAGAA CATGATTACA GGGCCTGCCC AGATGGANGG E. coli G. max Blb AAGAA TATGATTACA GGACCTGCTC AGATGGANGG E. coli G. max Blb AAGAA TATGATTACA GGACCTGCTC AGATGGANGG S. cerevisiaeb AAGAA TATGATTACA GGACCTGCTC AAATGGANGG A. thalianab AAGAA TATGATTACA GGACCTGCTC AAATGGANGG Selected sequence for AAAAA TATGATTACT GAGCTGCTC AAATGGANGG Selected sequence for AAAAA TATGATTACT GAGCTGCTC AAATGGANGG Selected sequence for Sequences or match capitals are indicated by lower-case letters. Mismatches for SEQ Indicate gaps in the sequences displayed. 30 "R" "Y" "M" "K" "W" "M" and "S" designate nucleotide positi stands for Inosine which is a nucleotide analog that can be the sequence refers to tuf (EF-1) gene. This sequence refers to tuf (AAAAA AAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	479	CCCGTCCGA CCTGTCGGA CCTGTTCGT CCTGTTCGT CCCCATCCG CCCATCCG TCCTATTAGA TCCTATTAGA TCCTATTAGA TCCTATTAGA TCCTATTAGA TCCTATTAGA		TATIAGR		which T; "W" to any	1 primers.
5 C. neoformansa AAGAA CATGATTACT GGTaCTtCTC AAGAA TATGATTACT GGTaCTtCTC AAGAA TATGATTACA GGGACATCT Humana AAGAA TATGATTACA GGGACATCT AAGAA CATGATTACA GGGCTGCCC G. max B2b AAGAA CATGATTACA GGGCTGCCC G. max B2b AAGAA CATGATTACA GGGCTGCCC G. max B2b AAAA CATGATTACA GGGCTGCCC G. max B2b AAAA CATGATTACA GGGCTGCCC S. aureofaciensc AAAAA CATGATTACA GGGCTGCCC S. aureofaciensc AAAAA TATGATTACA GGAGCTGCCC AAAAA TATGATTACA GGAGCTGCCCC AAAAA TATGATTACA GGAGCTGCCCCCCCCCC					cereví tch tl 3 <u>0</u> ID	sition or G c an bir	lected
5 C. neoformansa AAGAA CATGATTACT GGTaCTtCTC AAGAA TATGATTACT GGTaCTtCTC AAGAA TATGATTACA GGGACATCT Humana AAGAA TATGATTACA GGGACATCT AAGAA CATGATTACA GGGCTGCCC G. max B2b AAGAA CATGATTACA GGGCTGCCC G. max B2b AAGAA CATGATTACA GGGCTGCCC G. max B2b AAAA CATGATTACA GGGCTGCCC G. max B2b AAAA CATGATTACA GGGCTGCCC S. aureofaciensc AAAAA CATGATTACA GGGCTGCCC S. aureofaciensc AAAAA TATGATTACA GGAGCTGCCC AAAAA TATGATTACA GGAGCTGCCCC AAAAA TATGATTACA GGAGCTGCCCCCCCCCC	9	SgetGAC' NgetGAC' SgetGAC' SATGGAC SATGGAC SATGGAC AATGGAT AATGGAT	RATGGA		nyces s or ma for S	ide pos cands f that c	•
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C. neoformansa AAGAA CATGAT S. cerevisiaea AAGAA TATGAT Humana 10 G. max B1b AAAAA CATGAT G. max B2b AAAAA CATGAT G. max B2b AAAAA CATGAT E. colic S. aureofaciens AAGAA TATGAN S. cerevisiaeb AAGAA TATGAN A. thalianab A. thalia		Gracet Gracrt Gracrt GGaCA GCGCTG GCGCCG GCGCCG GAGCCG GAGCCG	GIGCI		·H 3	ayed. late nu or C; tide a	gene. organ rom ba mpleme
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		C. ne Numar G. m. G. m. F. c. c. F. c. c. A. c. c.	Sele	Sele	The capi are		σουσ
<i>- 1 - 3</i>		5 10 10 15			25	30	35

tuf from eukaryotic sequencing primers of selection Annex VII:Strategy for the (EF-1) sequences.

SEQ ID Accession NO.: #:	665 x00779 - D64080	- M29934	_ U81803	- M92073	- D14542 - D14542	x03558	_ U72244	- M64333	- AJ224150	- AJ224153	_ U42189	- L76077	- AF054510		558 560
314	TITI AGAGA TITCATCAAG AACATGATTA CTGG	WITCATCAAG AATATGATCA CTGG	CTTCATCAAG AACATGATCA CCGG	TITCATIAAG AACAIGAITA CIGG		CITCAICAAG AACAIGAICA CIGG	CHILATCAAA AACATGATTA CAGG	CGCGA CITCAICAAG AACAICAICA COCCIII	HICHTARA AATATGATTA CTGG	TIII PARACA III AND AND AND AND AND COGG	TGTT AAGGA III CAILLE AACAGATTA CCGG	TITICGIGA LITERIA DACATGATCA CGGG	TACGCGTGGG TCTTCGCGA CITCAICA ABCATGATCA CCGG		GA YITCAIYAAR AAYAIGAIYA C
179 286	TT AGAGA T	TGCTCGTGA	TOIL:::AGAGA	TCTTAGAGA I	TCCTCGCGA C	TCCTCGTGA C	TCTTAGAGA		TATTCGIGA		T.T. AMGGA	T.T			
154			TACGCTTGGG	TACGCITIGGG		ATGCGTGGG		ACGCGTGGG	TATGCTTGGG		TACGCATGGG	Trecricang Tacgeeriege	TCTTTCAAG	GG TICTITCAAG TACGCTTGGG II	TCITTYAAR TAYGCITGGG T
.T	s cerevisiae G			C. neoformans			Suracum							Y. lipolytica	Selected sequences for amplification primers
v	,	. ~		10	•	- '		15	2			27	20	i	;

sequences. Mismatches for SEQ ID no. 558 and 560 are indicated by lower-case letters. Mismatches for SEQ ID NG. The sequence numbering refers to the Saccharomyces cerevisiae tuf (EF-1) gene fragment (SEQ ID NO. Nucleotides in capitals are identical to the selected sequences SEQ ID NOs. 558, 560 or 653, or match 653 are indicated by underlined nucleotides. Dots indicate gaps in the sequences displayed. 30

stands for C or T; "M" stands for A or C; "K" stands for G or T; "W" stands for A or T; "S" stands for C or G. "I" "R" "Y" "M" "K" "W" and "S" designate nucleotide positions which are degenerated. "R" stands for A or G; stands for inosine which is a nucleotide analog that can bind to any of the four nucleotides A, C, G or T. 35

tuf from sequencing primers eukaryotic selection of Annex VII:Strategy for the (EF-1) sequences (continued)

D À	665 X00779 - D64080 - M29934 - U81803 - U81803 - D14342 - U14100 - X03558 - A07224150 - AJ224153 - AF054510	654	655 559
1304	GTTTACAA GATCGCTGCT ATTGGTACGACATG AGACAAACTG TCGCTGTCGG TGT GTGTACAA GATTGGCGGT ATTGGTACGATATG AGACAGACTG TCGCTGTCGG TAT GTGTACAA GATCGGTGGT ATTGGTACGATATG AGACAGACGG TTGCTGTCG TGT GTTTACAA GATCTCGGTG ATTGGAACGATATG AGACAGACG TTGCTGTTGG TGT GTTTACAA GATTCTCGG ATTGGAACGATATG AGACAAACG TTGCTGTTGG AGT GTGTACAA AATTGGTGGT ATTGGCACGATATG AGACAAACG TTGCTGTTGG TGT GTGTACAA AATTGGTGGT ATTGGACCGATATG AGACAAACG TTGCTGTTGG TGT GTGTACAA AATTGGTGGT ATTGGACCGATATG AGACAAACAG TTGCTGTTGG CAT GTTTACAA AATTGGTGGT ATTGGTACGATATG AGACAAACAG TTGCTGTTGG TAT GTATACAA AATTGGTGGT ATTGGTACGATATG AGACAAACAA TTGCTGTTGG TAT GTATACAA AATTGGTACGATATG AGACAAACAA TTGCTGTTGG TAT GTATACAA AATTGGTACGATATG AGACAAACAA TTGCTGTTGG TAT GTATACAA AATTGGTACGACATG CGCCAAACGA TTGCTGTTGG TAT GTGTTACAA AATTGGTACGACAATG CGCCCAAACGA TTGCTGTTGG TAT GTGTTACAA AATTGGTACGACAATG CGCCCAAACGA TTGCTGTTGG TAT GTGTTACAA AATTGGTACGACAATG CGCCCAAACGA TTGCTGTTGG TAT GTGTTACAAA AATTGGTACGACAATG CGCCCAAACGA TTGCTGTTGG TAT GTGTTACAAA AATTGGTACGACAATG CGCCCAAACGG TCGCTGTTGG TAT GTGTTACAAA CAATCGGTACGACAATG CGCCCAAACGG TCGCTGTTGG TAT GTGTTACAAA CAATCGGTACGACAATG CGCCCAAACGG TCGCTGTTGG TAT GTGTTACAAA CAATGGTACGACAATG CGCCCAAACGG TCGCTGTTGG TAT GTGTTACAAA CAATCGGTACGACAATG CGCCCAAACGG TCGCTGTTGG TGT GTGTTACAAA CAATCGGTAC ATTGGTACGACAATG CGCCCAAACGG TCGCTGTTGG TGT GTGTTACAAA CAATCGGTACGACAATG CGCCCAACGG TCGCTGTTGG TGT GTGTACAAACG TCGCTGTTGG TGT GTGTACAAACG TCGCTGTTGG TGT GTGTACAAACG TCGCTGTCGG TGT GTGTACAAAACG TCGCTGTCGG TGT GTGTACAAACG TCGCTGTCGC TGT GTGTACAAACG TCGCTGTCGG TGT GTGTACAACG TCGCTCGTCGC TGT GTGTACAAACG TCGCTCGTCGC TGT GTG		ATG MGICARACIR TYGCYGTCGG
776 1276	GGTACGACATG GGTACGATATG GGTACGATATG GGGACGATATG GGGACGATATG GGGACGATATG GGGACGATATG GGGACGATATG GGGACGATATG GGGACGATATG GGGACGATATG GGTACGATATG GGGACGATATG	99,1	
	GTTTACAA GATCGGTGGT ATT GTGTACAA GATTGGCGGT ATT GTTTACAA GATCGGTGGT ATT GTTTACAA GATCGGTGGT ATT GTCTACAA GATCTGGGG ATT GTCTACAA AATTCGTGGT ATT GTGTACAA AATTGGTGGT ATT GTGTACAA AATTGGTGGT ATT GTTTACAA AATTGGTGGT ATT GTGTACAA GATCGGTGGT ATT GTGTACAA GATCGGTGGT ATT GTGTACAA GATCGGTGGT ATT GTGTACAA GATCGGTGGT ATT	TACAA GAICGGIGGI ATYGG	TACAA RATYKGIGGT ATYGG
751	GTTTAC GTGTAC GTTTAC GTTTAC GTTTAC GTGTAC GTGTAC	TACAA	TAC
	S. cerevisiae B. hominis C. albicans C. neoformans E. histolytica G. lamblia H. capsulatum Human L. braziliensis O. volvulus P. berghei P. knowlesi S. pombe T. cruzi	Y. lipolytica Selected sequence for amplification primer	Selected sequences ror amplification primers ^a
5	9 S 8 275	25	

Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated cerevisiae tuf (EF-1) gene fragment (SEQ ID NO. by lower-case letters. "~" indicate incomplete sequence data. Dots indicate gaps in the sequences displayed. The sequence numbering refers to the Saccharomyces 30

stands for C or T; "M" stands for Å or C; "K" stands for G or T; "W" stands for A or T; "S" stands for C or G. stands for inosine which is a nucleotide analog that can bind to any of the four nucleotides A, C, G or T. "R" "Y" "M" "K" "W" and "S" designate nucleotide positions which are degenerated. "R" stands for A or G; 35

a This sequences are the reverse-complement of the selected primers.

s agalactiae- <i>specific</i>
Streptococcus sequences.
for the selection of Streptococcus ation primers from tuf sequences.
x VIII: Strategy
Annex

			· ON OIL	יייים יוסירא מסירע
		334 517 542	SEQ ID NO	
		W PRESCUE AND A PROPERTY OF THE PROPERTY OF TH	207	ı
'	S. adalactiae	TACIC PLANTAGE AND STATE OF THE	208	
•	S. adalactiae	TACTG ACAMACCTTT ACIIGGGC MACGENGER	209	1
	99-40-01-01-0	SCHOOL AMOUNTED	010	•
	S. agaractae	TACTG ACAAACCITT ACITGGAC AACGIIGGIG	077	
	S. agalactiae	かからしてはなりなったのかのか	211	ı
	S. anginosus	COCCUPACION TO THE TOTAL PROPERTY OF THE PROPE	221	1
2	v	TACTG ACAAACCATT GOILnone andraced	212	ı
•	; c	ACAAACCATT GCTT GGAC AACGIIGGIG	223	•
		CACTG ACAMACCATT GCTTAGAt AAtGTAGGTG ICCTTCTTC	777	,
	S. gordonii	ACAAGCCGGT CCTT GGAt AAtGITGGTG TICICCTICG	57.7	•
	S. mutans	ははして ままっているべるしゃ	145	
	S. pneumoniae	ひはい 単手して 単手しているがって	227	•
15	S. sanguinis	ACAMOCCALL GOLLS COMPONIA PACCAMOGRAGA	228	1
	S. sobrinus	COCCATACTE ACARCCATA GOLL	16	t
	D CODACÍA	ACGGCGCTT CCTGCGAC AACGALGGGAC ACCAGGGCGCGCGCGCGCGCGCGCGCGCGCGCGC	,	P33165
		CGCGATOTIG ATAAACCIII CIIGIGAC AACGIAGGIG 1911911100		700104
	B. fragilis	子であるのである。		FOTCES
	B, subtilis	CGCGACACIG MAMMACCAIL COIC CODO DACEGAGGTO	662	ı
20	ن	CGTGAGACGG ACAAGCCA11 CC1C1:100:10 11:00 00:00 0	22	1
Ì		OHOUSE STORY OF THE STORY OF TH	78	1
	יייי ביוסייים ביוס	CGTGGGALTG ACAAGCGTT CCTGTGAG AACGTAGGIG ILCIGGIGG	, r	
	E. coli	CONTRACTOR ACTION CONTRACTOR CONTRACTOR INCINCTICES	135	
77	G. vaginalis	CACCALCIA ACAACCATT CATOTGAC	179	1
	S. aureus			
3				
	Selected sequence for	A THEOLOGICAL CECAFACECE AND	549	
	species-specific primer			
			,	
		C AACGTTGGTG TTCTTCTTC	550	
30) species-specific primer			
	The sequence numbering ref	Streptococcus agalactiae tuf gene fragment (SEQ ID NO.	209). Nucleot: lower-case let	209). Nucleotides in capitais are lower-case letters. Dots indicate
	identical to the selected sequences			

"R" "Y" "M" "K" "W" and "S" designate nucleotide positions which are degenerated. "R" stands for A or G; "Y" stands for C or T; "W" stands for A or C; "K" stands for G or T; "W" stands for A or C; "K" stands for inosine which is a nucleotide analog that can bind to any of the four nucleotides A, C, G or T. gaps in the sequences displayed. 35

 40^{-6} The SEQ ID NO. refers to previous patent publication WO98/20157. b This sequence is the reverse-complement of the selected primer.

Strategy for the selection of Streptococcus agalactiae-specific hybridization probes from tuf sequences. Ann x IX:

SEQ ID NO.: Accession #: 206 209 144* 207 210 228 221 212 213 214 215 216 217 219 220 222 223 224 P33170 - Genome project 226 146* 227 228 229 229 229	582 583
431 433 GGTACTOT TABABETHAL CACCAAGTEG AANTCOTTGG TATTAAACA CHARTCOLAN AACCAGTTGT TA GGTACTOT TEAGTCAAC CACCAAGTEG AANTCOTTGG TATTAAACAA CATATCOLAN AACCAGTTGT TA GGTACTOT TCCTGTCAAC CACCAAGTTG AANTCOTTGG TATTAAACAA CATATCOLAN AACCAGTTGT TA GGTACTOT TCCTGTCAAC CACCAAGTTG AANTCOTTGG TATTAAACAA CATATCOLAN AACCAGTTGT TA GGTACTOT TCCTGTCAAC CACCAAGTTG AANTCOTTGG TATTAAACAA CATATCCAAN AACCAGTTGT TA GGTACTOT TEAGTCAAC CACCAAGTTG AANTCOTTGG TATCOGTAAC CACAATCCAAN AACCAGTTGT TA GGTACTOT TEAGTCAAC CACCAAATCG AANTCOTTGG TATCOGTAAC CACAATCCAAN AACCAGTTGT TA GGTACTOT TEAGTCAAC CACCAAATCG AANTCOTTGG TATCOAGAGA CAAATCCAAN AACCAGTTGT TA GGTACTOT TEAGTCAAC CACCAAATCG AANTCOTTGG TATCOAAGAA CAAATCCAAN AACCAGTTGT TA GGTACTOT TEAGTCAAC CACCAAATCG AANTCOTTGG TATCOAAGAA CAAATCCAAN AACCAGTTGT TA GGTACTOT TEAGTCAAC CACCAAATCG AANTCOTTGG TATCOAAGAA CAAATCCAAN AACCAGTTGT TA GGTACTOT TEAGTCAAC CACCAAATCG AANTCOTTGG TATCAAAGAA CAAACCAAN AACCAGTTGT TA GGTACTT TEAGTCAAC CACCAAATCG AANTCOTTGG TATCAAAGAA CAAACCAAA AACCAGTTGT TA GGTACTT TEAGTCAAC CACCAAATCG AANTCOTTGG TATCAAAACAAA AAAACAAAA AACCAGTTGT TA GGTACTT TEAGTCAAC CACCAAATCG AANTCOTTGG TATCAAACAAAAA AACCAGTTGT TA GGTACTT TEAGTCAAC CACCAAATCG AANTCOTTG	actgt tcgtgtcaac gacgaagttg aaa cgttgg tattaaagaa gatatccaaa aagcagttg
5 s. acidominimus s. agalactiae s. agalactiae s. agalactiae s. agalactiae s. agalactiae s. agalactiae s. anginosus s. anginosus s. anginosus s. anginosus s. cricetus s. cricetus s. cricetus s. cricetus s. dysgalactiae s. dysgalactiae s. gordonii s. macacae s. gordonii s. macacae s. gordonii s. macacae s. preumoniae s. progenes s. preumoniae s. progenes s. pretis s. sanguinis s. sanguinis s. sanguinis s. suberis s. suberis s. uberis s. uberis s. uberis	Sel spe
\$ 01 \$1 02 \$7 277	, 4

The sequence numbering refers to the *Streptococcus agalactiae tuf* gene fragment (SEQ ID NO. 209). Nucleotides in capitals are identical to the sequences or match those sequences. Mismatches are indicated by lower-case letters. Dots indicate gaps in the sequences displayed.

45

The SEQ ID NO. refers to previous patent publication WO98/20157. These sequences are the reverse-complement of the selected probes.

fic	SEQ ID 399	380 379 381 381 382 382 382 247 248 292 293 293 324 310 627	625 626
agalactiae- <i>specific</i>		GTTCCTT ATTAGCACCT TACTTAAAAG GTGGTAAAG GTTCCTT GTTAGCACCT TACTTAAAAG GTGGTAAAG GTTCCTT GTTAGCACCT TACTTAAAAG GTGGTAAAG GTTCCTT TATAGCACCT TACTTAAAAG GTGGTAAAG GTTCCTT TATAGCACCT TACTTAAAG GTGGTAAAG GTTCCTT ATTAGCCCCT TACTTAAAG GTGGTAAAG GTTCCTT ATTAGCCCCT TACTTAAAG GTGGTAAAG GTTCCTT ATTAGCCCCT TACTTAAAG GTGGTAAAG GTACTTT ATTAGCCCCT TACTTAAAG GTGGTAAAG GTACTTT ATTAGCTCT TACTTAAAAG GTGGTAAAG GTACTTT ATTAGCTCT TACTTAAAAG GTGGTAAAG GTACTTT ATTAGCTCT TACTTAAAAG GTGGTAAAG GTACCTT ATTAGCTCT TACTTAAAAG GTGGTAAAG GTACCTT ATTAGCACCT TACTTAAAAG GTGGTAAAG GTACCTT ATTAGCACCT TACTTAAAAG GTGGTAAAG GTACCTT ATTAGCACCT TACTTAAAAG GTGGTAAAAG GTACCTT ATTAGCACCT TACTTAAAAG GTGGTAAAA	AITAGCACCI TACTIAAAAG GTGGIA
of Streptococcus sequences.	234 368	TGTCTTCAAC GTTCCTT GGTCTTCAAC GTTCCTT GGTCTTCAAC GTTCCTT TGTTTTAAC GTTCCTT	ggata ctitgggicg igicticaac g
Δ	203	TAAGGATA C TAAGGATA C TAAGGATA C TAAGGATA C TAAGGATA C TAAAGGAA C	GGATA
X: Strategy for the selection amplification primers from atpD	99	TT GATTGTCTAT AAAATGGGG ATAAGTCACA TT GATTGTCTAT AAAATGGGG ATAAGTCACA TT CATTGTCTAT AAAATGGGG ATAAGTCACA TT GATTGTCTAT AAAATGGGG ATAAGTCACA TT GATTGTTTAT AAAATGGCG ATAAGTCACA TT GATTGTTTA AAAATGGCG ATAAGTCACA TT GATTGTTAT AAAAATGGCG AAAAGTCAA CT tGTCGTTAT AAAAATGGCG AAAAGACCCA TT GATGTTAT AAAAATGGCG AAAAGAGC TT GATGTTAT AAAAATGGCG AAAAAAAGC TT GATGTTTAT AAAAATGGCG AAAAAAAC TT GATGTTTAT AAAAATGGCG AAAAAAAC TT GATGTTTAT AAAAATGGCG AAAAAAAC TT GATGTTTAT AAAAATGGCG AAAAAAAAC TT GATGTTTAT AAAAATGGCG AAAAAAAAC TT GATGTTTAT AAAAAAGGCG AAAAAAAC TT GATGTTTAT AAAAAAGGCG AAAAAAACC TT AAATGGTCTAT AAAAAAGGCG AAAAAAACC TT AAATGGTCATAAAAAGAACAAC TA CAAAGAACAAC TA CATGGTTATA AAAAAACCCC AAAAAAACCC TT AAATGGCC AAAAAACCC TT AAAAAAACCCC AAAAAACCC TT AAAAAAACCCC AAAAAAACCC TT AAAAAAACCCC AAAAAAACCCC AAAAACCC TT AAAAAAAAACCCC AAAAAAACCCC AAAACCCC TT AAAAAAACCCC AAAAAAACCCC AAAACCCC TT AAAAAAAAACCCC AAAAAACCCC AAAACCCC TT AAAAAAAAAA	Selected sequences for species-specific primers ^y
Ann x X:		S. agalactiae S. agalactiae S. agalactiae S. agalactiae S. agalactiae S. agalactiae S. bovis S. bovis S. proumoniae S. progenes S. anginosus S. anginis S. mutans B. anthracis B. anthracis E. faecium E. faecium E. faecium E. faecium E. faecium E. faecium S. aureus S. aureus S. aureus S. aureus S. apidermidi Selected seque for species-s	
	5	01 51 02 52 08 278	35

The sequence numbering refers to the Streptococcus agalactiae tuf gene fragment (SEQ ID NO. 380). Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated by lower-case letters. Dots indicate gaps in the sequences displayed.

40 **d***** These sequences were obtained from Genbank and have accession #: a=AB009314, d=AF001955, e=U31170, and f=V00311.

** These sequences were obtained from genome sequencing projects.

** These sequences are the reverse-complement of the selected primers.

ic amplification la dubliniensis-	491 SEQ ID Accession NO.: #:	T 624 T 409	TTG T 410 -	∺ [⊷	€~ (TTG T 414 -	→ E	.	TTG T 421 -	Ę.	carc c 623 -	∺ F	· Æ	ပ	TTAT 447 -	• €		H.	577 578	s are identical to the ches for SEQ ID NO. 578	के ल	
ndida albicans/dubliniensis-specificic hybridization probe and Candidatuf sequences.	428 460	CACCAACTC AAATCCGGIA AAGTTACTGG TAAGACCTTG	AAATCCGGTA AAGTTACTGG	CACCAACTC AAATCCGGTA AAGTTACTGG TAAGACCTTG	C AAAICCGGIA AGGITACIGG	C AAATCCGGTA	CACCAACTC AAATCCGGTA AGGITACIGG TAAGACCTIG CACCAACGC AAGGCtGGTG tcGTcAagGG TAAGACCTTG	AAggoteera Agtocacoe	CACCAACGC AAggCtGGTA ccGTCAAgGG TAAGACCTTG	AAGTCYGGTA AGtccACcGG	AAgTCtGGTg tttccAagGG	AACCAACTT AAAGCtGGTA AGGTTACCGG TAAGACCTTG	AGGICACTGG	AAggatGGCA	AAAGCtGGTg	CACCAACAC AAGGCGGGG tGGTGAAGGG TAAGACTGTT		AICCGGIA AAGTIACIGG IAAGACCI	CCA (C. albicans) .CCA (C. dubliniensis)	(SEQ ID NO. 408). Nucleotides in capitals armare indicated by lower-case letters. Mismatches	splayed. d. "R" stands for A or G; "Y" stands for C " stands for inosine which is a nucleotide	:/20157, SEQ ID NOS. 11-12)
the selection of Car lida albicans-specif: idization probe from	368 403	AAAGACTGCAACATGA TTGAACCATC	CAACATGA TTGAACCATC	AAAGACTGCAACATGA TTGAACCATC		AAAGACTGCAACATGA TTGAAGCTTC	GITACAACCC AAAGACTGCAACAIGA ITGAAGCtIC CAC	LAAGACTGCAACATGA TTGAGGCTTC	AAAGATGCAACATGA TTGAAGCGAC	GTIACAACCC AAAGACTGCAACATGA TIGAAGCAIC CA	CAAGGCTGCAACATGT TGGAGGAGG	tAAAgCTGCAAtATGA TTGAACCATC	: taacgergcaacarda rreaagetre	CARCATGC TOGACCARD	AAAAACTGTAACATGA TTGAACCATC	GITACAACCC AAAGACTGCAACATGA ITGAAGCTAC UA GITTCAACCC CAAGACCGTAACATGA ITGAGCGGAC CA	TTACAACCC AAAGA		CAIGA ITGAACCAIC CACCA (C. CAIGA ITGAAGCIIC CACCA (C.	da albicans tuf gene fragment	equences. Mismatches for SEQ ID NO. Of any interesticles. Dots indicate gaps in the sequences displayed. te nucleotide positions which are degenerated. "R" stands for A or T; "S" stands for C or G. "I" stands	, G or T. ribed in a previous patent (publication WO98/20157, ement of the selected primer.
Annex XI: Strategy for primers, Canon specific hybi	337	CGTC AAGAAGGITG	C. albicans CGTC AAGAAGGTTG G	albicans CGTC AAGAAGGTTG	C. albicans CGTC AAGAAGGTTG G	dubliniensis CGTC AAGAAGGTTG	dubliniensis CGTC AAGAAGGTTG		kefyr CATC AAGAAGGICG	CATC AAGAAGGTTG	C. lusitaniae CGIC AMERAGETTE G	is CGTC AAGAAGGTIG	CGTC AAGAAGGTTG	aigatus CATC AAGAAGGICG	AAGAAAGTTG	iae TATC AAGAAGGTTG	Selected sequence for species-specific amplification primer [®] C AAGAAGGTTG GTTACAACCC	Selected sequence for species-specific amplification primer ^{4,b}	Selected sequences for species-specific hybridization probes	The sequence numbering refers to t	selected sequences or match those sequences. Mismatches for are indicated by underlined nucleotides. Dots indicate gaps "R" "M" "R" "M" and "S" designate nucleotide positions on a cream stands for G or T; "W" stands for G or T; "W" stands for A or T; "S" st	to any of the four nucleotides A, C, G or T. • C. albicans primers have been described in by This sequence is the reverse-complement of
	د			10			1.6	C			20	3		2	ج 79		30	35	40	?	45	

575 707

TIACCA GAAGGIACIG AAAIGGIIA TIACCA GAAGGIACIG AAAIGGIWA

c amplification
Staphylococcus-specific
of
the selection tuf sequences.
Strategy for primers from
XII:
Annex

						** 40.00000
			310 340 652	789	SEQ ID NO.:	
•	L		A CACTIACCA TOAR CONTINUE CANATICANAG CACTIACCA GAAGGIACTG AAATGGIAAT	TAAT GC	179	,
•	. s.	aureus	TOBACCTOCT CAAATCAAAG	35 ~~~	176	
	s.	aureus	CAGGCGGGG BORNOGEGGG GRAND GRA	TAAT GC	177	
	s.	aureus	CAGGCCGTGT TGAACGAGGA CAANACAAAAAAAAAAAAAAAAAAAAAAAA	_	180	•
	s.	aureus aureus	CAGGCCGTGT TGAACGTGGT CAAALCAAAGCACLIACGT GTTACGTACGG		181	1
•	ŝ	auricularis	CAGGCGIGIT IGAACGIGGI CAAALCAAAGCACCC	TTAT GC	182	•
=	0 s.	capitis capitis	CAGGCCGTGT TGAACGTGGT CAAATCAAACAACLIACCA SAAGGTACTG		183	ı
			CIGGROGICE TEAGCETCER CANGILLANGOCOCERTAINCE	TTAT GC	184	
	s.		TGAGGGGG CAMALCANAGOACCAANGG CHICCONG		185	
	s.	epidermidis	TEARCETEST CANALCASS DACTTACCA GAAGGTACAG	TTAT GC	141	ı
-	s.	epidermidis	CAGCCCGIGI ISANCGIGGI CAMMINANGCI INCOMING CONTROLL CONTROLL CANGGIRGING	TTAT GC	186	•
<u> </u>	ر ج	haemolyticus	CAGGCCGIGI IGANCGIGG CANANCADAG ANCHINGCA		188	•
	'n	haemolycicus	COGGETOR DANGER CRANCE CANADADA COCCUE	~~ ~~~	189	
	s.		CAGGCCGIGI ICANCGIGGI CANACADADO DACTIBOCA	TAAT GC	191	•
	s.		CAGGCCGIGI IGAACGIGGI CAAAICAAAGAACTTACCA GAAGGIACIG	TAAT GC	193	•
Č	ري دي	hominis	CAGGGGG MONACGEGG CAMPACANACCANACCANACCANACCANACCANACCANAC	~~ ~~~	194	•
2	o S	hominis	CAGGCCGIGI ISAACGIGGI CAMAICANNO:	TAAT GC	195	•
	'n.	hominis	CAGGCCGTGT TGAACGTGGT CAMMICANAG		196	1
	S.	hominis	CAGGCGIGI IGANGGIGGI CANNAGONO DADGATAGON	TTAT GC	197	•
	ŝ	lugdunensis	CAGGCCGTGT TEAACGTGGT CANALCAMAC		198	•
	'n.	saprophyticus	CAGGCCGTGT TGAACGTGGT CAAATCAAAG	TTAT GC	199	•
રી 28	S.	saprophyticus	CAGGCCGTGT TGAACGTGGT CAAALCAAAGAACTTACGT GESCCTTGCC		200	•
80	s.	saprophyticus	CAGGCCGIGIT ICAACGIGGI CAAAICAAAGAACIIACCA CAICCIIICAAACAAAAAAAAAAAAA		201	•
	s.	sciuri sciuri	CAGGCCGIGI IGAACGIGGI CAMAICACIG:GGCZZIGGG GIAAGGTAGG		187	1
	ŝ	warneri	CAGGCCGIGI IGAACGIGGI CAMICANGCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	~~ ~~~	192	1
•	s,	warneri	CAGGCCGIGI LGANCGIGGI CANNICATORIO	STIAT GC	202	•
~	30 S.	warneri	CAGGCCGTGT TGAACGTGGT CAAALAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA		•	299104
	æ	subtilis	CIGGCCGIGI BEAACGCGG LAAGICAAAGChect.coch Calcock		78	•
	ы	E. coli	BGAACGOGGI BECATICALAGGAACIYCCY GAAGCGGCGGG		138	•
	T.	L. monocytogenes	A CIGGECGIGI IGAACGIGGA CAAGILAAAGACBGILCUA GAAGEIACIG AMALES			
m	S Sel	Selected sequence for denus-specific primer	GGCCGIGI IGAACGIGGI CAAATCA		553	
	; ,					

The sequence numbering refers to the *Staphylococcus aureus tuf* gene fragment (SEQ ID NO. 179). Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated by lower-case letters. "~" indicate incomplete sequence data. Dots indicate gaps in the sequences displayed.

"R" "Y" "M" "K" "W" and "S" designate nucleotide positions which are degenerated. "R" stands for B or G; "Y" stands for C or T; "M" stands A or C; "K" stands for G or T; "W" stands for A or T; "S" stands for C or G. "I" stands for inosine which is a nucleotide analog that can bipd to any of the four nucleotides A, C, G or T.

45

The SEQ ID NO. refers to previous patent publication WO98/20157. These sequences are the reverse-complement of the selected primers.

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genus-specific primers^b Selected sequences for

280

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Annex XIII: Strategy for the selection of the Staphylococcus-specific hybridization probe from tuf sequences.

•		400 425 SEQ ID NO.:	Accession #:
	S. aureus	G TTGAAATGTT CCGTAAATTA TTAGA 179	-
10	S. aureus	G TTGAAATGTT CCGTAAATTA TTAGA 176	. -
	S. aureus	G TTGAAATGTT CCGTAAATTA TTAGA 177	-
	S. aureus	G TTGAAATGTT CCGTAAATTA TTAGA 178	_
	S. aureus aureus	G TTGAAATGTT CCGTAAATTA TTAGA 180	
	S. auricularis	G TAGAAATGTT CCGTAAATTA TTAGA 181	-
15	S. capitis capitis	G TAGAAATGTT CCGTAAATTA TTAGA 182	_
15	M. caseolyticus	G TAGAAATGTT CCGTAAATTA TTAGA 183	-
	S. cohnii	G TAGAAATGTT CCGTAAATTA TTAGA 184	-
	S. epidermidis	G TAGAAATGTT CCGTAAATTA TTAGA 185	-
	S. haemolyticus	G TAGAAATGTT CCGTAAATTA TTAGA 186	_
20	S. haemolyticus	G TAGAAATGTT CCGTAAATTA TTAGA 189	-
	S. haemolyticus	G TAGAAATGTT CCGTAAATTA TTAGA 190	
	S. haemolyticus	G TAGAAATGTT CCGTAAATTA TTAGA 188	-
	S. hominis	G TAGAAATGTT CCGTAAATTA TTAGA 196	-
	S. hominis	G TAGAAATGTT CCGTAAATTA TTAGA 194	-
25	S. hominis hominis	G TAGAAATGTT CCGTAAATTA TTAGA 191	-
	S. hominis	G TAGAAATGTT CCGTAAATTA TTAGA 193	-
	S. hominis	G TAGAAATGTT CCGTAAATTA TTAGA 195	-
	S. lugdunensis	G TAGAAATGTT CCGTAAATTA TTAGA 197	-
	S. saprophyticus	G TAGAAATGTT CCGTAAATTA TTAGA 198	-
30	S. saprophyticus	G TAGAAATGTT CCGTAAATTA TTAGA 200	_
	S. saprophyticus	G TAGAAATGTT CCGTAAATTA TTAGA 199	-
	S. sciuri sciuri	G TTGAAATGTT CCGTAAATTA TTAGA 201	_
	S. warneri	G TAGAAATGTT CCGTAAGTTA TTAGA 187	-
	S. warneri	G TAGAAATGTT CCGTAAGTTA TTAGA 192	-
35	S. warneri	G TAGAAATGTT CCGTAAGTTA TTAGA 202	•
	S. warneri	G TAGAAATGTT CCGTAAGTTA TTAGA 203	- 299104
	B. subtilis	G TTGAAATGTT CCGTAAgcTt cTTGA	299104
	E. coli	G TTGAAATGTT CCGcAAACTG CTGGA 78	_
	L. monocytogenes	G TAGAAATGTT CCGTAAATTA CTAGA 138°	_
40			
	Selected sequence for		
	genus-specific hybridi		
	zation probe	GAAATGTT CCGTAAATTA TT 605	

The sequence numbering refers to the *Staphylococcus aureus tuf* gene fragment (SEQ ID NO. 179). Nucleotides in capitals are identical to the selected sequence or match that sequence. Mismatches are indicated by lower-case letters.

^{50 *} The SEQ ID NO. refers to previous patent publication WO98/20157.

Annex XIV: Strat gy for the s lection of Staphylococcus saprophyticus-sp cific and of Staphylococcus haemolyticus-specific hybridization probes from tuf sequences.

5

								SEQ ID
10		339					383	NO.:
10	S. aureus	AG 1	T tGGTGAAGA	AgTtGAAATC	ATCGGTTTAC	ATGACACATC	TAA	179
	S. aureus	AG S	TEGGTGAAGA	AgTtGAAATC	ATCGGTtTaC	ATGACACATC	TAA	176
	S. aureus	AG '	TEGGTGAAGA	AGTEGAAATC	ATCGGTtTaC	ATGACACATC	TAA	177
	S. aureus	AG '	TEGGTGAAGA	AGTEGAAATC	ATCGGTTTAC	ATGACACATC	TAA	178
15	S. aureus aureus	AG '	TEGGTGAAGA	AgTtGAAATC	ATCGGTTTAC	ATGACACATC	TAA	180
13	S. auricularis	AG '	TCGGTGAAGA	AGTTGAAATC	ATCGGTATga	AaGACggTTC	AAA	181
	S. capitis capitis	AG '	TEGGTGAAGA	AgTtGAAATC	ATCGGTATCC	Acgaractic	TAA	182
	M. caseolyticus	AG '	TEGGTGAAGA	AgTtGAAATC	ATTGGTtTaa	cTGAagaacC	AAA	183
	S. cohnii	AG '	TCGGTGAAGA	AGTEGAAATC	ATCGGTATgC	AaGAagaTTC	CAA	184
20	S. epidermidis	AG '	TEGGTGAAGA	AGTEGAAATC	ATCGGTATgC	Acgaractic	TAA	185
20	S. haemolyticus	AG '	TEGGTGAAGA	AGTEGAAATC	ATTGGTATCC	ATGACACTTC	TAA	186
	S. haemolyticus	AG	TEGGTGAAGA	AGTEGAAATC	ATTGGTATCC	ATGACACTTC	TAA	189
	S. haemolyticus	ΔG	TEGGTGAAGA	AGTEGAAATC	ATTGGTATCC	ATGACACTTC	TAA	190
	S. haemolyticus	AG	TEGGTGAAGA	AgTtGAAATt	ATTGGTATCa	Aagaaacttc	TAA	188
25	S. hominis	AG	TEGGTGAAGA	AGTEGAAATE	ATTGGTATCA	AAGAAACTTC	TAA	194
	S. hominis hominis	AG	TtGGTGAAGA	AgTtGAAATt	ATTGGTATCA	AAGAAACTTC	TAA	191
	S. hominis	AG	TtGGTGAAGA	AgTtGAAATt	ATTGGTATCA	AAGAAACTTC	TAA	193
	S. hominis	AG	TEGGTGAAGA	AgTtGAAATt	ATTGGTATCa	AAGAAACTTC	TAA	195
	S. hominis	AG	TtGGTGAAGA	AgTtGAAATt	ATTGGTATCa	AAGATACTTC	TAA	196 197
30	S. lugdunensis	AG	TCGGTGAAGA	AgTtGAAATt	ATTGGTATCC	ACGATACTAC	TAA	— - -
	S. saprophyticus	AG	TCGGTGAAGA	AATCGAAATC	ATCGGTATGC	AaGAagaaTC	CAA	
	S. saprophyticus	AG	TCGGTGAAGA	AATCGAAATC	ATCGGTATGC	AAGAAGAATC	CAA	
	S. saprophyticus	AG	TCGGTGAAGA	AATCGAAATC	ATCGGTATGC	ABGABGBBTC	CAA	
	S. sciuri sciuri	TG	TtGGTGAAGA	AgTtGAAATC	ATCGGTTTAA	CTGAAGAATC	TWW.	
35	S. warneri	AG	TtGGTGAAGA	AgTtGAAATC	ATCGGTTTAC	ATGACACTIC	. ተማህ	
	S. warneri	AG	TEGGTGAAGA	AgTtGAAATC	ATCGGTTTAC	ATGACACTIC	י עמעי	
	S. warneri	AG	TtGGTGAAGA	AgTtGAAATC	ATCGGTCTAC	ATGACACTIC	י ארשי	
	S. warneri	AG	TtGGTGAAGA	AgTtGAAATC	ATCGGTETAC	ATGACACTIC	. 1777	
•	B. subtilis	AG	TCGGTGACGA	AgTtGAAATC	ATCGGTCTCC	. AAGAAGAGGG	CAN	•
40	E. coli	AG	TtGGTGAAGA	AgTtGAAATC	griggiarca	AAGAGACICO	. 222	138 ^b
	L. monocytogenes	AG	TtGGTGAcGA	AgTaGAAgTt	ATEGGTATES	, AaGAagaaag	, AA	150
	Selected sequences for							
	species-specific						•	500
45	hybridization probes		CGGTGAAGA	AATCGAAATC	A (S. sapi	rophyticus)	_	599
	-		(S. h	emolyticus)	ATTGGTATCO	C ATGACACTTO		594

The sequence numbering refers to the Staphylococcus aureus tuf gene fragment (SEQ ID NO. 179). Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated by lower-case letters.

^{*} This sequence was obtained from Genbank accession #299104.

b The SEQ ID NO. refers to previous patent publication WO98/20157.

Annex XV: Strategy for the selection of Staphylococcus aureus-specific and of Staphylococcus epidermidis-specific hybridization probes from tuf sequences.

5

```
SEQ ID
                                                    547
                                                          592
                                                                                   617
                                                                                         NO.:
                           521
                           TACACCACA TACTGAATTC AAAGCAG...TTCTTCtCa AACTATCGtC CACAATT
                                                                                          179
10
     S. aureus
                           TACACCACA TACTGAATTC AAAGCAG...TTCTTCtC- -----
                                                                                          178
     S. aureus
                           TACACCACA TACTGAATTC AAAGCAG...TTCTTCtCa AACTATCGtC CACAATT
                                                                                          176
     S. aureus
                           TACACCACA TACTGAATTC AAAGCAG...TTCTTCtCa AACTATCGtC CACAATT
                                                                                          177
     S. aureus
                           TACACCACA TACTGAATTC AAAGCAG...TTCTTCtCa AACTATCGtC CACAATT
                                                                                          180
     S. aureus aureus
                           TACACCACA CACTAAATTC ACtGCAG...TTCTTCtCT AACTACCGtC CACAATT
                                                                                          181
15
     S. auricularis
                           CACACCACA CACTEAATTC AAAGCGG...TTCTTCAGT AACTACCGCC CACAATT
     S. capitis capitis
                           TACECCACA TACTARATTC ARAGCTG...TTCTTCACT ARCTACCGCC CECAGTT
                                                                                          183
     M. caseolyticus
                                                                                          184
                           TACACCACA CACABACTTE AAAGCGG...TTCTTCAgT AACTATCGCC CACAATT
     S. cohnii
                           TACACCACA CACABAATTC ARAGCTG...TTCTTCACT ARCTATCGCC CACAATT
                                                                                           185
     S. epidermidis
                           CACACCECA CACAGAATTE AAAGCAG...TTCTTCACA AACTATCGEC CACAATT
                                                                                          186
20
     S. haemolyticus
                           CACACCECA CACABARTE AAAGCAG...TECTTCACA AACTATCGEC CACAATE
                                                                                          189
     S. haemolyticus
                           CACACCECA CACBBAATTE ARAGCAG...TTCTTCACB ARCTATCGEC CACAATT
                                                                                           190
     S. haemolyticus
                           TACACCECA CACBBARTE ARAGCAG...TTCTTCACT ARCTATCGEC CACAATT
                                                                                           188
     S. haemolyticus
                           CACACCECA CACABARTEC ARAGCAG...TTCTTCACT ARCTATCGEC CACAATT
                                                                                           195
     S. hominis
                           TACACCECA CACABAATTC AAAGCAG...TTCTTCACT AACTATCGEC CACAATT
                                                                                           196
25
     S. hominis
                           TACACCECA CACABAATTC AAAGCAG...TTCTTCECT AACTATCGEC CACAATT
                                                                                           191
     S. hominis hominis
                           TACACCECA CACBBAATTC AAAGCAG...TTCTTCECT AACTATCGEC CACAATT
     S. hominis
                            TACACCECA CACBBAATTC AAAGCAG...TTCTTCECT AACTATCGEC CACAATT
                                                                                           194
     S. hominis
                           TACACCECA CACTAATTE AAAGCTG...TTCTTCECA AACTACCGCC CACAATT
                                                                                           197
     S. lugdunensis
                           TACACCACA TACABAATTC AAAGCGG...TTCTTCACT AACTACCGCC CACAATT
                                                                                           198
30
     S. saprophyticus
                           TACACCACA TACBRAATTC AAAGCGG...TTCTTCACT AACTACCGCC CACAATT
                                                                                           199
     S. saprophyticus
                            TACACCACA TACABAATTC AAAGCGG...TTCTTCACT AACTACCGCC CACAATT
                                                                                           200
     S. saprophyticus
                            CACACCECA CACTARATTC ARAGCTG...TTCTTCACA ARCTACCGCC CACAATT
     S. sciuri sciuri
                            TACACCACA TACABAATTC AAAGCGG...-----
                                                                                           192
    S. warneri
                                                                                           187
                            TACACCACA TACABAATTC AAAGCGG...TTCTTCAgT AACTACCGCC CACAATT
     S. warneri
35
                            TACACCACA TACABARTTC AAAGCGG...TTCTTCAGT AACTACCGCC CACAATT
                                                                                           202
     S. warneri
                            TACACCACA TACABAATTC AAAGCGG...TTCTTCAGT AACTACCGCC CACAATT
                                                                                           203
     S. warneri
                            CACTCCACA CAGCARATTC ARAGCTG...TTCTTCTCT ARCTACCGTC CTCAGTT
     B. subtilis
                                                                                           78
                            CAAGCCGCA CACCBAGTTC GAATCTG...TTCTTCABB GGCTACCGTC CGCAGTT
     E. coli
                                                                                           138<sup>b</sup>
                            TACECCACA CACTAACTTC AAAGCTG...TTCTTCAAC AACTACCGCC CACAATT
40
     L. monocytogenes
     Selected sequences
     for species-specific
     hybridization
                                                                                           585
                               ACCACA TACTGAATTC AAAG (S. aureus)
45
     probes
                                                                                           593
                                             (S. epidermidis) TTCACT AACTATCGCC CACA
```

The sequence numbering refers to the Staphylococcus aureus tuf gene fragment (SEQ ID NO. 179). Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated by lower-case letters. "~" indicate incomplete sequence data. Dots indicate gaps in the sequences displayed.

This sequence was obtained from Genbank accession #Z99104.

The SEQ ID NO. refers to previous patent publication WO98/20157.

Annex XVI: Strategy for the selection of the Staphylococcus hominis-specific hybridization probe from tuf sequences.

S. aureus

S. aureus

ATC ATCGGTTTAC AtGACACATC TAA
ATC ATCGGTTTAC ATCACACATC TAA
ATC ATCGGTTATC ACGARACTTC TAA
ATC ATCGGTTATC ACGARACTTC TAA
ATC ATCGGTTATC AAGAACACTC TAA
ATC ATTGGTATCC ATCACACTTC TAA
ATC ATTGGTATCC ATCACACTTC TAA
ATC ATTGGTATCA AAGAACACTTC TAA
ATC ATTGGTATCA AAGAACCTTC TAA
ATC ATTGGTATCA AAGAACCTTC TAA
ATC ATTGGTATCA AAGAACCTTC TAA
ATC ATTGGTATCA AAGAACCTTC TAA
ATC ATTGGTATCA AAGAAACCTTC TAA
ATC ATTGGTATCA AAGAAACCTTC TAA
ATT ATTGGTATCA AAGAAACCTTC TAA
ATT ATTGGTATCA AAGAAACCTTC TAA
ATT ATTGGTATCA AAGAAACCTTC TAA
ATC ATCGGTATGC AAGAAACCTTC TAA
ATC ATCGGTTATCA AAGAACCTTC TAA
ATC ATCGGTTATCA AAGAAACCTTC TAA
ATC ATCGGTTATCA AAGAAACCTTC TAA
ATC ATCGGTTATCA AAGAAACCTTC TAA
ATC ATCGGTTATCA AAGAAACCTTC TAA
ATC ATCGGTTATCA AAGAACCTTC TAA
ATC ATCGGTTATCA AAGAAACCTTC TAA
ATC ATCGGTTATCA AAGAAACCTTC TAA
ATC ATCGGTTATCA AAGAAACCTTC TAA
ATC ATCGGTTATC SEQ ID NO.: -78 138^b Selected sequence for species-specific hybridization probe ATTGGTATCA AAGAAACTTC

- 5

The sequence numbering refers to the *Staphylococcus aureus* tuf gene fragment (SEQ ID NO. 179). Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated by lower-case letters. Dots indicate gaps in the sequences displayed.

a This sequence was obtained from Genbank accession #Z99104.

b The SEQ ID NO. refers to previous patent publication WO98/20157.

amplification Enterococcus-specific the of selection primers from tuf sequences. the for Strategy Ann x XVII:

			i i i i i i i i i i i i i i i i i i i					
		270	298	556	582	SEQ ID NO.:	Accession #:	
3	E. avium	TAGAATTAAT GG	GGCTGCTGTT GACGAATAT.	TGAA GATATCCAAC	GTGGACAAGT ATT	131ª	•	
	E. casseliflavus	TGGAATTAAT GG	GGCTGCAGTT GACGAATAC.	TGAA GACATCCAAC	GTGGACAAGT ATT	58	ı	
	E. cecorum	TAGAATTAAT GG	GGCTGCAGTT GACGAATAC.	TGAA GATATCCAAC	GTGGtCAAGT ATT	59	•	
	E. dispar	TAGAALTAAL GO	GOCTGCAGTT GACGAATAT	TGAA GATATCCAAC	GIGGECAAGT ATT	09	ı	•
	E. durans	TTGAATTAAT GG	GGCTGCAGTT GACGAATAT	TGAA GACATCCAAC	GIGGACAAGI TIT	61	•	
9	E. flavescens	TGGAALTAAT GG	GGCTGCAGTT GACGAATAC	TGAA GACATCCAAC	GIGGACAAGI AIT	65		
	E. faecium	TTGAATTAAT GO	GOCTGCAGTT GACGAATAC.	TGAA GACATCCAAC	GIGGACAAGI TIT	608	•	
	S. faecalis	TAGAATTAAT GG	GGCTGCAGTT GACGAATAT.	TGAA GATATCGAAC	GTGGACAAGT ATT	607	,	
	E. gallinarum	TGGAATTBAT GG	GGCTGCAGTT GACGAATAC	TGAA GACATCCAAC	GIGGACAAGI ATT	609	ı	
	E. hirae	TTCAATTGAT GG	GGCTGCAGTT GACGAATAT.	TGAA GACATCCAAC	GTGGACAAGT TTT	29	•	
15	E. mundtii	TTGAATTGAT GG	GGCTGCAGTT GACGAATAT	TGAA GACATCCAAC	GIGGECAAGI TIT	89		
	E. pseudoavium	TAGAATTAAT GS	GSCTGCTGTT GACGAATAC TGAA	TGAA GACATCCAAC	GIGGACAAGI ATT	69	•	
	E. raffinosus	TAGAATTAAT GG	GGCTGCTGTT GATGAATAC TGAA	TGAA GACATCCAAC	GIGGACAAGI AIT	70	•	
	E. saccharolyticus	TCGAATTAAT GG	GGCTGCAGTT GACGAATAT.	TGAA GACATCCAAC	GTGGACAAGT ATT	71		
	E. solitarius	TGGACTTAAT GG	GGATGCAGTT GATGACTAC	TGAE GATATCGAAC	GTGGtCAAGT ATT	72	•	
20	E. coli	TGGAACTEGG to	togetteerg dartetTAY	TGAA GABATCGAAC	greetcager ACT	7.8	1	
	B. cepacia		odacdCgorg GACaogTAC.	TGAA GACGTGGAGC	Greechger TCT	16	•	
	B. fragilis		GGaaGCTGTT GATACTTGG.,	GAAC GAAATCAAAC	Greetatger TCT	ť	M22247	
	B. subtilis	TCGAACTEAT GG	GGATGCGGTT GATGAGTAC	TGAA GABATCCAAC	GTGGtCAAGT ACT	1	299104	
	C. diphtheriae	TCGACCTCAT GC	GcaggCTtgc KATGAtTCC	. CGAA GACGITGAGC	Greecager Ter	662	•	
25	C. trachomatis	GAGAGCTAAT GC	GCBBGCCGTC GATGATAATGAAC	GAAC GATGTGGAAG	GaGGAAtgGT TGT	22	•	
	G. vaginalis	AGGAACTCAT Ga	GAAGGCTGTT GACGAGTAC.	TACC GACGTtGAGC	GIGGLCAGGT TGT	135	•	
		TAGAATTART GG	GGaaGCTGTa GATactTAC	TGAA GACGTACAAC	Gredtcaagr ATT	179	1	
	S. pneumoniae	TGGAATTGAT GA	GARCACAGTT GATGAGTAT	TGAE GABATCGAAC	GIGGACAAGI TAT	145	ſ	
	A. adiacens	TAGAATTAAT GG	GGCTGCTGTT GACGAATAC	TGAA AACATCGAAC	GIGGACAAGI TCT	118		
30	G. haemolysans		GGARRCAGTT GACGAATACTGAA	TGAA GACATCGAAC	GIGGACAAGI TIT	87	•	
	G. morbillorum		GGRARCAGIT GACGAGTAC TGAA	TGAA GATATCGAAC	Grogacaagt ttt	88	ı	
	Selected seguence for							
7	amplification primer	AATTAAT GG	AATTAAT GGCTGCWGTT GAYGAA			1137		
3	Selected sequence for amplification orimer			A GAYATCSAAC GTGGACAAGT	GTGGACAAGT	1136		

The sequence numbering refers to the Enterococcus durans tuf gene fragment (SEQ ID NO. 61). Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated by lower-case letters. Dots indicate gaps in the sequences displayed.

"Y" "W" and "S" designate nucleotide positions which are degenerated. "Y" stands for C or T; "W" stands for A or T; "S" stands for C or G. "I" stands for inosine which is a nucleotide analog that can bind to any of the four nucleotides A, C, G or T.

The SEQ ID NO. refers to previous patent publication W098/20157.
 This sequence is the reverse-complement of the selected primer.

hybridization probe, of the Enterococcus faecium-specific hybridization faecalis-specific probe and of the Enterococcus cassellflavus-flavescens-gallinarum group-Enterococcus specific hybridization probe from tuf sequences. the of selection the for Strategy Ann x XVIII:

Accession #:		•	•		•	•		ι		,			•		1	•	•	1	•	Z99104	.•		M22247	1				4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4
SEO ID NO.:	131	58	89	09	61	62	809	65	609	67	89	69	70	71	72	662	135	16	179		145	7.8	•	22		1174	602	7711
675	. GATACLITGE IACGIGGIGT	GGTGCATTGC	GGTGCATTAL	GGTGCATTat	GGTGCtTTaC	GGTGCtTTat	GGTGCTTTAC		GGTGCATTGC	GGTGCtTTaC	GGTGCgTTaC				GGTACLTTGE	GGTCLGCTtC		GGTatccTGC	GGTGCATTAL	GOTGCCCTCC	というないのできる		Serence of	GGICLBIIGC	2212822828		cfum)	T GGTGCAFTGC TACGTGG
	OPC RAPA	GGTATCGCT	GGIATIGGI	GGTATCCA1		COMPARTOR AND	GGIAIIAAAAAAAA		GGTATTGCT	GGTATCGCA	GGTATCGCA.	GGTATCGCT	GGTATTGCT		. :	GOTATCCGC		GGILLE GCGC		TGRAATCATC GGTCTBCALCALA	TGARATCATC GGICLICAN	GGTATGAAA	TGARATOGET GGTATCARACGTA	collarcte dortraddrtera	rcagtrggrc @Gretragaculr		18) aagt tgaagtroff gotatt (B. faecium)	
		GTGACGAAGT	GTGACGAAGT	GTGACGAAGT	GTGACGAAGT	GTGACGEEGE	GTGACGAAGT	GTGACGAAGT	GTGACGAAGT	GTGATGAGG	GTGACGCCCT	GTGACGECAT	GTGACGAAGT	GTGACGAAGT	GTGACGEEGT	GCGATGAAGT	ACGAGGACGI	ACACCCCAGE		GTGAAGAAGT	GTTBBBGTCG GTGACGAAGT	GITABAGTCA ACGACGARAT	GTGAAGAAGT	GTGATGAAAT	CCGATAAAGT		GTTCGC (E. faecalis)	
	395	GITTON ACGIGGACAN (GTTGA ACGTGGacAA	GTTGA ACGTGGacAA	GTTGA ACGTGGBCAA	GTTOA ACGTGGTGAA	GITTON ACGIGGOOM	GITTON ACGIGGGCAN	GTTON ACGIGGACAN	GTTON ACGTGGaCAN	GTTOR ACGTGGacAA	GTTGA ACGTGGacAA	GTTGA ACGTGGacAA	GTTGA ACGTGGBGAA		-	GTTGA SCGTGGTAAS	GTCGA gCGcGGcatc	GITGA ACGIGGICAA	GTAGA ACGCGGGCAA	ATCOM CCOTOCOTOCO	CTAGA ACGCGGTAto	ATCON AACTGGTGtt	ATTON SCOTGGGGtt	for	GA ACGTGGTGAA	86
		Bi.;334 &	a caccolfflavis	F CENTER TO THE	E dispar	F durans	E faccalis	E. faecium	E. flavescens	E. gallinarum	E. hirae	E. mundtil	E. pseudoavium	R raffinosus	E. saccharolyticus	E. solitarius	C. diphtheriae	G vacinalis	D CODOCIA	6. surella	E subrilis		S. pneumoniae	E. COLL	C. trachomatis	TOT Residence Particles	species-specific or	group-specific hybridization probes
S				2	2				15					00,1	2 28				30	3				ç	₹			32

The sequence numbering refers to the Enterococcus faecium tuf gene fragments (SEQ ID NO. 608). Nucleotides in capitals are identical to the sequences or match those sequences. Mismatches are indicated by lower-case letters. Dots indicate gaps in the sequences displayed. • The SEQ ID NO. refers to previous patent publication W098/20157.

the selection of primers for the identification of platelets contaminants from tuf sequences. Strategy for Annex XIX:

z99104 #:		
Accession 299104		
SEQ ID NO.: 7 7 7 78 100 103 153 209 140* 159 118 224 - 146* 235	636	637
TRAACTC AGTTCTACTT CCGTACACT GACGTAAC TRAACTC AGTTCTACTT CCGTACACT GACGTAAC CAAACTC AGTTCTACTT CCGTACACT GACGTAAC CAAACTC AGTTCTACTT CCGTACACT GACGTGAC CAAACTC AGTTCTACTT CCGTACACT GACGTGAC CAAACTC AGTTCTACTT CCGTACACT GACGTGAC CAAACTC AGTTCTACTT CCGTACACT GACGTGAC CAAACTC AGTTCTACTT CCGTACTACT GACGTGAC CAAACTC AGTTCTATTT CCGTACTACT GACGTGAC TAAATTC AATTCTATTT CCGTACTACT GACGTGAC TAAACTC AGTTCTACTT CCGTACTACT TAAACTC AGTTCTACTT CCGTACTACT TAAACTC AGTTCTACTT CCGTACTACT TAAACTC AGTTCTACTT CCGTACAACT TAAACTC AGTTCTACTT TAAACTC AGTTCTACTT TAAACTC AGTTCTACTT TAAACTC AGTTCTACTT TAAACTC AGTTCTACTT TAAACTC AGTTCTACTT TAAACT TAA	YAA	TTCTAYTT CCGTACIACT GACGT
GTA ACTGGTGTAG AGATGTTCCG GTT ACAGGTGTTG AAATGTTCCG TGT ACTGGCGTTG AAATGTTCCG TGT ACTGGCGTTG AAATGTTCCG TGT ACTGGCGTTG AAATGTTCCG TGT ACTGGCGTTG AAATGTTCCG GTT ACTGGCGTTG AAATGTTCCG GTT ACTGGCGTTG AAATGTTCCG GTT ACTGGCGTTG AAATGTTCCG GTT ACTGGTGTTG AAATGTTCCG	ACTGGYGTTG ALATGTTCCG YAA	
B. cereus B. subtilis E. cloacae E. coli K. oxytoca K. pneumoniae P. aeruginosa S. agalactiae S. aureus S. choleraesuis S. epidermidis S. marcescens S. marcescens S. marchae S. salivarius S. sanguinis Y. enterocolitica	Selected sequence for amplification primer	Selected sequence for amplification primer ^b
5 10 20	25	

to in selected sequences or match those sequences. Mismatches are indicated by lower-case letters. Dots indicate gaps The sequence numbering refers to the E. coli tuf gene fragment (SEQ ID NO. 78). Nucleotides in capitals are identical sequences displayed. the

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"R" "Y" "M" "K" "W" and "S" designate nucleotide positions which are degenerated. "R" stands for A or G; "Y" stands for C or T; "M" stands for A or T; "W" stands for I or G or T; "W" stands for I or T. 35

[•] The SEQ ID NO. refers to previous patent publication W098/20157.
• This sequence is the reverse-complement of the selected primer.

atpD Strategy for the selection of the universal amplification primers from sednences. Annex XX:

: Accession #: X76875 Z73419 	Genome project	X76877 U32730 U43738 V00267 M22247	
SEQ ID NO.:291 380 -366	243 264 284 351 317 357	393 670	562 564 565 563
	ACCICCAGG ACCICCAGG ACCICCAGGA ACCACCAGGA ACCACCAGGA ACCACCGGGA ACCACCGGGA ACCACCGGGA	CGCCGTATG CGCCGTATG CGCCGTATG CGCCGTATG CGCCGTATG TCACGTATG	C ARATGRAYGA RCCICCIGGI GYIMGIATG TAYGGIC ARATGAAYGA RCCICCIGGI AA ATH CCITCIGCIG TIGGITAYCA RCC ATG CCITCIGCIG TIGGITAYCA RCC
	S. aureus A. baumannii N. gonorrhoeae C. freundii E. cloacae E. coli S. typhimurium K. pneumoniae	S. marcescens Y. enterocolitica B. cepacia H. influenzae M. pneumoniae H. pylori B. fragilis	Selected sequences for universal primers Selected sequences for universal primers ^a
5 01	15	දි දැ 288	30

The sequence numbering refers to the *Escherichia coli atpD* gene fragment (SEQ ID NO. 669). Nucleotides in capitals are identical to the sequences or match those sequences. Mismatches for SEQ ID NOs. 562 and 565 are indicated by lower-case letters. Mismatches for SEQ ID NOs. 562 in the sequences displayed.

[&]quot;R" "Y" "M" "K" "W" and "S" letters designate nucleotide positions which are degenerated. "R" stands for A or G; "Y" stands for C or T; "M" stands for A or T; "H" stands for A, C or T; "S" stands for C or G. "I" stands for inosine which is a nucleotide analog that can bind to any of the four nucleotides A, C, G or T. 8

a These sequences are the reverse-complement of the selected primers.

Sp cific and ubiquitous primers fr nucleic acid Annex XXI: amplification (recA sequences).

		Originatin	g DNA fragmen
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
	Universal primers (recA)		
919	5'-GGI CCI GAR TCI TMI GGI AAR AC	918 ^a	437-459
920b	5'-TCI CCV ATI TCI CCI TCI AIY TC	918ª	701-723
921	5'-TIY RTI GAY GCI GAR CAI GC	918 ^a	515-534
922b	5'-TAR AAY TTI ARI GCI YKI CCI CC	918 ^a	872-894
	Sequencing primers (recA)		
1605	5'-ATY ATY GAA RTI TAY GCI CC	1704 ^a	220-239
1606	5'-CCR AAC ATI AYI CCI ACT TTT TC	1704 ^a	628-650
	Universal primers (rad51)		
935	5'-GGI AAR WSI CAR YTI TGY CAY AC	939 ^a	568-590
936p	5'-TCI SIY TCI GGI ARR CAI GG	939a	1126-114
	Universal primers (dmc1)		
937	5'-ATI ACI GAR GYI TTY GGI GAR TT	940ª	1038-106
938p	5'-CYI GTI GYI SWI GCR TGI GC	940 ^a	1554-157

a Sequences from databases.

35

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex XXII: Specific and ubiquitous primers for nucleic acid amplification (speA sequences).

5			Originatin	g DNA fragment
	SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
)	Bacterial s	pecies: Streptococcus pyogenes		
	994	5'-TGG ACT AAC AAT CTC GCA AGA GG	993a	60-82
	995b	5'-ACA TTC TCG TGA GTA ACA GGG T	993 ^a	173-194
j	996	5'-ACA AAT CAT GAA GGG AAT CAT TTA G	993 a	400-424
	997b	5'-CTA ATT CTT GAG CAG TTA CCA TT	993a	504-526
	998	5'-GGA GGG GTA ACA AAT CAT GAA GG	993a	391-413
)	997b	5'-CTA ATT CTT GAG CAG TTA CCA TT	993 ^a	504-526

a Sequence from databases.

²⁵ b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

pyogenes-sp cific Streptococcus amplification primers from speA sequences. selection of the First strategy for Ann x XXIII:

SEQ ID NO.:	66 	1 1 1 1 1			
adratGTGAtCCT.GT egtTCAtGAG adratGTGAtCCT.GT egtTCAtGAG gGTATGTGACCT.GT TACTCACGAG GGTATGTGACCT.GT TACTCACGAG GGTATGTGACCCT.GT TACTCACGAG	GGACTAACAA TCTCGCAAGA GGTATGTGACCCT.GT TACTCACGAG	TCTCGCAAGA GGTATGTGACCCT.GT TACTCACGAG TCTCGCAAGA GGTATGTGACCCT.GT TACTCACGAG TCTCGCAAGA GGTATGTGACCCT.GT TACTCACGAG TCTCGCAAGA GGTATGTGACCCT.GT TACTCACGAG TCTCGCAAGA GGTATGTGACCCT.GT TACTCACGAG	GGACT GGACT GGACT GGACT GGACT	GGTATGTGACCCT.GT TACTCACGAG GGTATGTGACCCT.GT TACTCACGAG aGGTAGTGACCCTGGT TACTCACGAG	ACCCT.GT TACTCACGAG AATGT
Accession # X61573 A AF029051 A X61571 A X61570 A X61570				spea X61564 spea X61565 spea AF055698 spea X03929° Selected sequence for species-specific primer	Selected sequence for species-specific primer
S spek spek spek spek	10 spea spea spea spea	15 spea spea spea spea	20 spea spea spea spea spea spea spea	spea spea spea spea 30 Sele	Sel 35 spe

The sequence numbering refers to the Streptococcus pyogenes speA gene fragment (SEQ ID NO. 993). Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated by lower-case letters. "~" indicate incomplet sequence data. Dots indicate gaps in the sequences displayed.

The extra G nucleotide introducing a gap in the sequence is probably a sequencing error.

This sequence is the reverse-complement of the selected primer.

pyogenes-sp cific selection of Streptococcus speA sequences. from the Second strategy for amplification primers Ann x XXIV:

	Accession # X61573 AF029051	TTTAGAAAAAAAATGGT AACTGCTCAA	SEQ ID NO.:
X61571 X61570		A ACAAATCATG AAGGGAATCA TTTAGAAAAAAAATGGT AACTGCTCAA A ACAAATCATG AAGGGAATCA TTTAGAAAAAAAATGGT AACTGCTCAA A ACAAATCATG AAGGGAATCA TTTAGAAAAAAAATGGT AACTGCTCAA	
X61568 X61569		CGGAGGGGTA ACAATCATG AAGGGAATCA	1 1
X61572 X61560		CGGAGGGGTA ACALATCATG AAGGGAATCA TITAGAAA AAAAATGGT	993
U40453 X61554		ACAMATCATG AAGGGAATCA TTTAGAAAAAAAATGGT AACTGCTCAA	, ,
X61557 X61559		AACTGCTCAA	1 1
X61558			i t
X61556 X61555		CGGAGGGGTA ACAAATCATG AAGGGAATCA TTTAGAAAAAAAATGGT	
X61560		AACTGCTCAA	•
x61566		CGGAGGGTA ACAAATCATG	
X61567		AACTGCTCAA	ı
x61563		COGREGEGTA ACAMATCATE	1 1
X61564		. AAAAATGGT AACTGCTCAA	1
X61565 AF055698		COGRAGOGIA ACAAATCATG AAGGGAATCA TTTAGAAAAAAAATGGT AACTGCTCAA	
X03929		TA COGROGOGTA ACADATCATO ARGGORATCA TTTAGAAAAAAAATGGT AACTGCTCAA GAATIAG.CI	
Selected sequences for species-specific prime	Selected sequences for species-specific primers	ggagggta acaaatcatg aagg	966
Selected sequence for species-specific primer*	ce for	AAIGGT AACTGCICAA GAATTAG	766

The sequence numbering refers to the Streptococcus pyogenes speA gene fragment (SEQ ID NO. 993). Dots indicate gaps in the sequences displayed.

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This sequence is the reverse-complement of the selected primer.

cific	SEQ ID	NO.:	211	212	217	1002	1448	985	145	215	982	200	227	225	146	231	229	224	226	222	214	607	176	7	78		(999	1001	1000
pyogenes-specific		647	CATCACACA CTAAATT			_	_			_	_	_	_	_	_	_	_	_	CATCCGCACA CTAAATT	-	CALCCACACA CTAAATT	ACTCCACACA CANANT	ACACCACACA CTGAATT	_	AAGCCGCACA CCAAGTT					AGTTCAATC AACCCACACA CTAA
Streptococcus es.		186 619		THE KUMMUTE TO	THE STREET OF TO	CHANGEACEACOCC	C AMGIICAMIC.	CC Agerrcant	CC AgGTTCAATC	CC Agentraarc	CASCAMONACIONAL CONTRACTOR CONTRA	THE STREET STREET	CTARCITUDE OC.	AGGTTCAATC	CO TOGETTCAATC	ა ლ :		د د		د	<u>်</u>	ر ا		ָרָ מַרְ		} •			GAG	AGTTCAATC
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for the				GTTGACGAtG	GTTGATGACG	GTTGAcGAtG	GTTGATGACG	GTTGATGAtG	GTAGACGACG	GTTGAcGACG	GTTGATGACG	GTTGATGACG	GTTGAcGAtG	GTTGAcGAtG	GTTGATGAtG	GTTGAcGAtG	GTTGAcGAtG	GTTGAcGAtG	GTTGACGAtG	GTTGATGATG	GTTGATGAtG	GTTGACGACG		GTTGACGAtG	GTAGATGACG	GTTGATGACG		TAPERTERCE		
Strategy for	amplification F		140	A AGTTGACTTG	A AGTTGACCTT	A AATTGACCTT	A AGTTGACCTT	A AGTTGACCTT	A AATTGACTT	A AGTIGACTI	A GATCGACTT	A GATCGACTT	A AGTTGACTT9	A AGTTGACTTG	A AGTTGACTT9	A AGTTGACTTG		A AGTTGACTT9	A AGTTGALTY						A ATBCGACATG	A ATGCGACaTG				۵
	ambi			anginosus	ž.	dysgalactiae	enes	ace action	is	pneumoniae	cristatus	to.	gorđonii	sanguinis	parasanguinis ,	salivarius	vestibularis		su	·-	ıcae	cricetus	faecalis	sna	sne	••		Selected sequences for	species-specilic primers	Selected sequence for species-specific primer ^b
Ann x XXV:				S S. andil	, v	S. dvsa			10 S. oralis	, c	•				S. para				620 S. mutans	S. ratti	S. macacae	S. cric	E. faec	25 S. aureus	B. cereus	E. coli			30 species	Selecte

The sequence numbering refers to the Streptococcus pyogenes tuf gene fragment (SEQ ID NO. 1002). Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated by lower-case letters. Dots indicate gaps in the sequences displayed.

The SEQ ID NO. refers to previous patent publication W098/20157.

⁴⁰ b This sequence is the reverse-complement of the selected primer.

Ann x	XXVI	Strategy	for	the	selection	stx,-specific	Strategy for the selection stx_i -specific amplification primers and	primers	and
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		q	hybridization probe.	S	SEQ ID
				175 391 421	NO.:
		Accession #	263 343		
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	SEX	M21534	ArgtttAtch gGaGCGTACaG	TITIO ACTUAL TATCATION TO	
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	, X	X81415	COAGGGCTTG ATGTTTATCA GGGCCGTACAG	TITIETH TACKET TO THE TOTAL TO	
00	, y ,	x81416	caagggertg Argettatca ggagggTACag	TTTGCACATA TACCALISAITCCA TGACAACGGA	
3	SFX	X81417	Argittatch gGaGCGTACaG	Trigcacata istantis anticca igacasegga	
	Stx	X81418	Argettatca gGaGCGTACaG	TITIGACAMA MATCACTIC GITTECA TGACAGGGA	
	SEX	E03962	ATGECTATER GGCGCGTACCG	TITACACATA TATCACACA GITTECA	•
	of X	E03959	gGCGCGTACCG	TIIACACATA TACCACACACACACACACACACACACACACAC	
بر 29	,	x07865	goccccTACCG	Track All taccacies control and the control of the	
	2,50	X10775	gaegeg TACCG	Triachchia Iaccadioorgania Trachchia	1077
	, tr	237725	goccccTACCG	TTTBCACATB TACCACIOGITCCCA TGBCBACGGA	,
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	stx;	AF043627	TGGATATA CGAGGGCTtG ATGTCTAtcA gGcGCGTACAG A		
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	Selec	Selected sequence for		Characteria and the conformation of the confor	1084
	hybri	hybridization probe			
4	Selec	Selected sequence for	иe	ACAT TGTCTGGTGA CAGTAGCTAT A	1080
	ildme	amplification primer"		the conitals are identical to the	

The sequence numbering refers to the Escherichia coli stx, gene fragment (SEQ ID NO. 1076). Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated by lower-case letters. Dots indicate gaps in the sequences displayed.
• This sequence is the reverse-complement of the selected primer.

1079

AATCAGCA ATGTGCTTCC G

1085

the selection of stx_2 -signobe.	the selection of probe.	for the selection of tion probe.	for the selection of tion probe.	the selection of probe.	pecific amplification primers and	
selection .	the selection probe.	for the selection tion probe.	for the selection tion probe.	for the selection tion probe.	of stx,-specific	
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				35Q 1D NO
		Accession #	2) O 11 A AGGETGAGE CECAGTGGC GTECTTAA AGGETGAGE GCGCCTGC	•
S	stxi	M19473	TOTICACION SOCIONA SOCIONA SOCIONA SOCIONA SOCIONA SOCIA SOC	•
	stxi	M16625	TOTLACTOTO ACACAAC ACTUGRISH CHARLES ALCOHOLD	
	X	M17358	CAAC ACTGGBTGAC CCCAGIGGGC BICCIIA NOSTTGBGTA	
	S X	236900	ACA CAAC ACTGGGTGAT CTCAGTGGGC GICCIIA ACATGGGGT GTGTCCTGCC	•
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9	יין יין	1000	metricont rottactors ACA CAAC ACTGGGTGAt ctcAgTGGGC gftcTTA A AGGLIGAGLA BIGICLISC	1
2	SCX	CANTE	markacom montacron aca Caac Acresicat ctcagregge greetla A Acgrigate greetigee	1
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	SCX	236899	TOTLACTOTO ACA CAAC ACTUGATOR COCCASTOBAC MACCATOR	1076
	Stx	236901	ACA CAAC ACTEGATGAC CCCAGIGGEC SICCLIA	1
2	S. C.	X61283	ACAAGGC ACTGTCTGAAACTGCTC CIGIGIAG COMMINGER ATGTGCTTCC	•
:	S. T. X.	1,11079	AGGC ACTGTCTGA AACTGCTC CTGTGTA O ACABACAGCA ATGTGCTTCC	•
	2 4	M2153d	CIGILIA G AGMAICAGO A MONOCOMO	•
	SCX2	*CCT7E	MINISTER TOTAL TOTAL TOTAL ACA TOTAL TOTAL TOTAL TOTAL CONTINE	ı
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	stx2	U72191	Trendential Terescient Achterials Actions and American Samuel Control of the Cont	•
20	Stx2	X81415	TICIGCOTT TOTCACTOR ACA	•
	Stx2	X81416	TOTCACTOTC ACATGGC ACTUICIDAACTGGTC CTGTTTD G	•
	Stx,	X81417	ACA. TGGC ACTGTCTAAAACTGCTC CHGATTAA.	•
	Stx	X81418	ACATGGC ACHGICHGAAACIGCIC CIGILIA G CGBATCAGCA	
	Stx	E03962	ACAAGGC ACTGICTEAAACTGCTC CIGIGIA	•
25	S. C.	E03959	C ACAAGGC ACTIGICIGA AACTIGITA CIGIGIA CONSTITUTO ATGREENTIC	•
15	Stx	X07865	ACAAGGC ACTGICTGAAACTGCTC CIGIGIAG COMMICAGGA	•
	y X	Y10775	ACAAGGC ACTGTCTGA AACTGCTC CIGIGIAN O COBBITCAGE	1077
	stx	237725	C ACA AGGC ACTGICTIGA AACTGCIC CIGIGIA G CGBATCAGCA	•
	stx	250754	ACAAGGC ACTGICTGA MACTGCTC CIGIGIO C COMMICAGEA	
Ç	S. T. X.	X67514	ACA AGG ACTGTCTGA AACTGCTC CIGIGIA C NOBBTCBGCA	•
2	S X	L11078	ACAAGGC ACTOTOTOTA AACTOCITC CIGIGIA G ACEBATOROR ATGROCITCO	
	, y	X65949	ACAAGGC ACTOTOTOTAAACTGCTC CTGTGTAG AGASTGCC TGTGTCCTCC	
	stx,	AF043627		
č	. !			1078
5	Selec	Selected sequence for amplification primer	AG INCIGCOINT TGICACTGIC	8/01

295

The sequence numbering refers to the Escherichia coli stx, gene fragment (SEQ ID NO. 1077). Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated by lower-case letters. Dots indicate gaps in the sequences displayed.

C ACTGTCTGA. .. AACTGCTC CTGT

45

Selected sequence for amplification primer

Selected sequence for hybridization probe

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primers	
amplification	
vanA-specific	
of	
selection	
the	•
For	aces
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Ann x XXVIII:	
Ann	

	SEQ ID NO.: 1139 1141 1051 1052 1053 1054 1055 1056 1057 1049 1050 1117	1089
van sequences.	GTCAAT AGCGCGGACG ANTTGGACTA CGT AGAGGTCTAG CCCGTGTGGA TATG GTCAAT AGCGCGGACG ANTTGGACTA CGT AGAGGTCTAG CCCGTGTGGA TATG GTCAAT AGCGCGGACG ANTTGGACTA CGT AGAGGTCTAG CCCGTGTGGA TATG GTCAAT AGCGCGACG ANTTGGACTA CGT AGAGGTCTAG CCCGTGTGGA TATG GTCAAT AGCGCGGACG ANTTGAACTA CGT AGAGGTCTAG CCCGTGTGGA TATG GTCAAT AGCGCGGAGG AATTGAACG TGC AGAGGGCTAG CCCGTGTGA TATG GTAAAC GGLACGGAAG AACTLAACGC TGC AGAGGGCTG CCCGTGTGA TCTT GTAAAC AGLACGGAAG AACTLAACGC TGC AGAGGGCTG CCCGTGTGA TCTT GTAAAC AGLACGGAAG AACTBAACGC TGC AGAGGGCTG CCCGTGTGA TCTT GTAAAC AGACGGAAG AACTBAACGC TGC AGAGGGCTG CCCGTGTGA TCTT GTAAAC AGACGGAAG AACTBAACGC TGC AGAGGGCTG CCCGTGTGA TCTT GTAAAC AGAC	GAGGICTAG CCCGTGTGGA T
Ann x XXVIII:	Accession # vanA X56895 vanA vanB U94526 vanB U94527 vanB U94529 vanB U94529 vanB U94529 vanB U94529 vanB U94529 vanB U94530 vanB U72704 vanB	Selected sequence for amplification primer
	\$ 01 15 00 30 . \$2 08	35

The sequence numbering refers to the Enterococcus faecium vanA gene fragment (SEQ ID NO. 1139). Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated by lower-case letters. Dots indicate gaps in the sequences displayed.

This sequence is the reverse-complement of the above selected primer.

from amplification primers vanB-specific of selection Strategy for the van sequences. Annex XXIX:

	SEQ ID NO.:	1139	1141	1051	1052	1001	1000	**************************************	1055	1056	1057	1049	1050	111./	ı	1	ı	•	1		t			•	1	1	i		1095		1096	tines in softhoothing in 1911
real pedicers.	495 608 633	Coreattosa toggeragae Astat ACG GASTCTTCG tattcatcag GAA		DOTABLE DOS CONTRACTOR DESCRIPTION OF THE PROPERTY OF THE PROP	COGCARGAC AATAL ACG GGALCALCOC CITECOCTOC	AATAT ACG GRATCITECG CAICLAICAG	AATAT ACG GAATCTTECG tATTCATCAG	CATECATCAG	. tcggcaagac aatatacg gaatcritcg tartcarcag	. tCgGCAaGAC AATATACG GAATCTTtCG tATtCATCAG	teggeragae Aatatacg gaarerteg tarteareag	LCGCCAAGAC AATAT ACG GAATCTTLCG LATLCATCAG	. tCgGCAAGAC AATATACG GAATCITtCG tATtCATCAG	GCGGCAGGAC AATATACG GTATCTTCCG CATCCATCAG	GCAGCAGGAC AATAT ACG GTATCTTCCG CATCCATCAG	GCGGCAGGAC AATATACG GTATCTTCCG CATCCATCAG	A GCGGCAGGAC AATAT ACG GTATCTTCCG CATCCATCAG	A GCGGCAGGAC AATATACG GTATCTTCCG CATCCATCAG	A GCGGCAGGAC AATATACG GTATCTTCCG CATCCATCAG	A GCGGCAGGAC AATATACG GTATCTTCCG CATCCATCAG	GCAGCAGGAC AATATACG GTATCTTCCG CATCCATCAG	A GCGGCAGGAC AATATATG GTATCTTCCG CATCCATCAG	CATCCATCAG	_	A GCAGCAGGAC AATAT ACG GTATCTTCCG CATCCATCAG	GAAGCAAGAA AATATACG GCTTTTTAA GATTCATCAG	AGCAATAGAC GAAGCTTCAA AATATATG GCTTTTTCGA CTATGAAGAG AAA		は、 これのひをしむれている。		GENERAL CATCCATCAG	6
A TTP A	4.0000000	ACCESSION # ACCESSION	CEODEA		vanA - A (vanA - A (- A	4				: <	. 4	1194526 C	1194527 C	1194528 C	1194529	1194530	283305	U81452	135369	U72704	1.06138		1100456		AF136925 A		Selected sequence for	ampintication primer	Selected sequence for	amplification primer
			n n	Š	Š	Š	Š	01		> :	> =	> 2	, <u>,</u>		> 2	> 2	> 2) 97	• >	. 2		25		•	د .		30	U)	10		35

The sequence numbering refers to the *Enterococcus faecium vanB* gene fragment (SEQ ID NO. 1117). Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated by lower-case letters. Dots indicate gaps in the sequences displayed.

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Strategy for the selection of vanC-specific amplification primers from vanC sequences. Annex XXX:

••																		dentical to
SEQ ID NO.:	1058	1059	1138	1060	1061	1062	1063	i	•	1064	1065	1066	ı		1101		7071	enitals are
1092	GT CGACGGTYTT TYTGATTITG AAGAGAAACGGGTC TGGCTCGAAT CGATTTTTTC GT	CT CGACGGTTTT TYTGATTTTG AAGAGAAACGGGTC TGGCTCGAAT CGATTTTTTC GT	CT CONCRETET TITGATITIG ANGAGAAACGGGIC IGGCICGAAI CGAITITITC GI	GT AGACGGCTTT TYCGATTTTG AAGAAAAAAGGTC TYGCTCGCAT CGACTTYTTT GT	GT AGACGGCTTT TICGATTTTG AAGAAAAAAGGTC TIGCICGCAT CGACTTTTTT GT	ITCGATITIG AAGAAAAAAAGGIC ITGCICGCAI CGACITITII GI	TICGALTITIG AAGAAAAAAAGGIC TIGCICGCAI CGACTITITIT GI	TTCGATTTTG AAGAAAA AAAGGTC TTGCTCGCAT CGACTTTTTT GT	TICGATITIG AAGADAAAAAGGIC TIGCICGCAI CGACTITITI GI	TTCGATITIG AAGAAAAAAAGGTC TTGCTCGCAT CGACTTITIT GT	TICGAITITIG AAGAAAAAAAGGIC IIGCICGCAI CGACITITITI GI	TTCGATTITIG AAGAAAAAAAGGAC TTGCTCGCAT CGACTTTTTT GT	micramming aagaaaa Aaaggic tigcicgcai cgactiffit GT				GGTC TRGCTCGMAT CGAYTTTT	and the section in conitals are identical to
1064	ACGGGTC	. ACGGGTC	. ACGGGTC	. AAAGGTC	. AAAGGTC	AAAGGTC	AAAGGTC	AAAGGTC	AAAGGTC	AAAGGTC	. AAAGGTC	AAAGGaC	AAAGGTC				GGTC	
957	AAGAGAA.	AAGAGAA.	AAGAGAA.	AAGAAAA.	AAGAAAA.	AAGAAAA.	AAGAAAA.	AAGAAA	AAGAAAA.	AAGAAAA.	AAGAAAA.	AAGAAAA.	AAGAAAA		AAGA			
	PTTGATTTTG	PTTGATTTG	FTTGATTTTG	TTCGATTTTG	TTCGATTTTG	TTCGATTTTG	TTCGATTTTG	TTCGATTTTG	TTCGATTTTG	TTCGATTTTG	TTCGATTTTG	TYCGATTTTG	للمدررية والملململين		TIYGATITIG AAGA			
•	CGACGGTTTT	CGACGGTTTT	CONCEGUTIT	AGACGGCTTT	AGACGGCTTT	GT AGACGGCTTT	GT AGACGGCTTT	GT AGACGGCTTT	GT AGACGGCTTT				With Cook of	GT AGACGGCTTT	GACGGYTTT			
929	<u>.</u>	; <u>f</u>	ָּלָ פֿל	5 6	5 5	g Fg	GT	GT	GT	G	ָּבָּ כֿל	ָל ל	5 6	5				
# מסיממים		i I		76167W		ı	1	L29638	129638			1		L29639	Selected sequence	for resistance primer	Selected sequence	IOF resistance primer
	5	Vanci	Vancı	vancı	Vance	vance	vanC2	vanC2	Chen		Valid.	vancs	vancs	vanC3	Select	for rea	Selecto	ior re
	•	n				٥	2					2				70		

298

i. 2 The sequence numbering refers to the vanCl gene fragment (SEQ ID NO. 1138). Nucleotides in capitals are identical selected sequences or match those sequences. Mismatches are indicated by lower-case letters. Dots indicate gaps sequence displayed.

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ullet This sequence is the reverse-complement of the selected sequence.

pneumoniae-specific SEQ ID NO.: amplification primers and hybridization probes from pbp1a sequences. Streptococcus of selection the for Strategy Annex XXXI:

NO.: NO.: 1014 1017 1007 1008 1009 1011 1012 1016 1013 1016	1130
GA. TATATO ATGACGGAA TGATGAAAAC CGT GA. TATATO ATGACGGAA TGATGAAAAC GGT GA. TATATO ATGACGGAA TGATGAAAAC AGT GA. TATATO ATGACGGAA TGATGAAAAC TGT GA. TATATO ATGACGGAA TGATGAAAAC TGT GA. TATATO ATGACGGACA TGATGAAAAC TGT GA. TATATO ATGACGGACA TGATGAAAAC AGT GA. TATATO ATGACGGAAA TGATGAAAAC TGT GA. TACATO ATGACGGAAA TGATGAAAAC TGT GA. TATATO ATGACGAAA TGATGAAAAC TGT	atg atgachgama tgatgaaaac
A TUGACTACC AAGCATECAG TATCCTAATG CLATTTCAAG TAATACAACC A TCGACTACC AAGTATCAG TACCCAAATG CCATTTCAAG TAACACAACC A TCGACTACC AAGCATGCAT TATCCAAATG CCATTTCAAG TAACACAACC A TCGACTATCC AAGCATGCAT TATGCAAACG CCATTTCAAG TAACACAACA A TCGACTATCC AAGCATGCAT TATGCAAACG CCATTTCAAG TAATACAACA A TCGACTATCC AAGCATGCAT TATGCAAACG CCATTTCAAG TAACACAACA A TCGACTATCC AAGCATGCAT TATGCAAACG CCATTTCAAG TAACACAACA A TCGACTATCC AAGCATGCAT TATGCAAACG CCATTTCAAG TAACACAACA A TCGACTATCC AAGCATGCAT TATGCAAACG CCATTTCAAG TAACACAACT A TCGACTATCC AAGCATGCAT TATGCAACG CCATTTCAAG TAACACAACT A TCGACTATCC AAGCATGCAT TATGCAAACG CCATTTCAAG TAACACAACT A TCGACTATCC AAGCATGCAT TATGCAACG CCATTCAAG TAACACAACT A TCGACTATCC AAGCATGCAT TATGCAACG CCATTCCAAG TAACACACT A TCGACTATCC AAGCATGCAT TATGCAACG CCATTCCAAG TAATACAACT A TCGACTATCC AAGCATGCAT TATGCAACG CCATTCCAAG TAATACAACT A TCGACTATC	GACTATCC AAGCATGCAT TATG
Accession # M90528 X67873 AB006868 AF046234 AB006873 AF139883 AF159448 X67867 Z49094 X67870 X67870 X67871	Selected sequences for amplification primers
pppla	Select 35 amplif

the The sequence numbering refers to the Streptococcus pneumoniae pbpla gene fragment (SEQ ID NO. 1004). Nucleotides in capitals are identical to selected sequences or match those sequences. Mismatches are indicated by lower- case letters. Dotes indicate gaps in the sequences displayed.

CAAACG CCATTTCAAG TAATACAAC

Selected sequence for hybridization probe

8

[&]quot;R" "V" "M" "K" "W" and "S" designate nucleotide positions which are degenerated. "R" stands for A or G; "Y" stands for C or T; "M" stands for A or T; "H" stands for A, C or T;"S" stands for C or G. "I" stands for inosine which is a nucleotide analog that can bind to any of the four nucleotides A, C, G or T.

Annex XXXI:	Stratedy	for	the	selection		of	Stre	treptococcus		umoniae	pneumoniae-specific	i C
	amplification		K	and	hybridization	dizat	ion	probes	Erom	from pbpla	sedanc s	Ø
	(continued)	3).										

	SEQ ID NO.:	•	•	•		1014	1017		1169	1004	1007	1008	1009	1011		1005	1015	1006	1012	1 1	1010	1	1013	1016		1018		1	1193		1131
(continued).	756 783 813 840	GCTGGTAR BACLGGTACH TCBARCTATAA ATACGGGTTA TGTAGCTCCG GACGAAA	BACAGGBACC TCTAACTATA.	BACAGGBACC TCTAACTATA.	GCAGGIAA AACAGGIACT ICTAACIATA A ACACIGGIIA CGIAGCICCA GAIGAAA	GCAQGIAA AACAGGIACT ICIAACIAIA A ACACIGGIIA CGIAGCICCA GAIGAAA	GCTGGTAA GACAGGTACT TCTAACTACAA ACACTGGCTA TGTAGCTCCA GATGAAA	GCAGGTAA GACAGGTACT TCTAACTATAA ACACTGGCTA CGTAGCTCCA GATGAAA	AACAGGAACC TCTAACTATAA ACACTGGCTA TGTAGCTCCA	GCTGGTAA AACAGGAACG TCTAACTATAA ACACTGGCTA TGTAGCTCCA GATGAAA	AACAGGAACC TCTAACTATAA ACACTGGCTA TGTAGCTCCA	BACAGGAACC TCTAACTATAA ACACTGGCTA TGTAGCTCCA	GCTGGTAA BACAGGBACC TCTAACTATAA ACACTGGCTA TGTAGCTCCA GATGAAA	GCTGGTAA AACAGGAACG TCTAACTATAA ACACTGGCTA TGTAGCTCCA GATGAAA	CCTGGTAR BACAGGBACG ICTARCTATAA ACACTGGCTA IGTAGCICCA GAIGAAA	GCTGGTAN GACAGGTACT TCTAACTACAA ACACTGGCTA TGTAGCTCCA GATGAAA	GACAGGIACT ICTAACTACAA ACACTGGCTA TGTAGCTCCA	GACAGGTACT TCTAACTACAA ACACTGGCTA TGTAGCTCCA	GACAGGTACT TCTAACTACAA ACACTGGCTA TGTAGCTCCA	GACAGGTACT TCTAACTACAA ACACTGGCTA TGTAGCTCCA	CGTAGCTCCA	GACAGGTACT TCTAACTATA.	GACAGGIACT TCTAACTATAA ACACTGGCTA CGTAGCTCCA	GCAGGIAA GACAGGIACI ICIAACIATAA ACACIGGCIA CGIAGCICCA GAIGAAA	GACAGGTACT TCTAACTATAA ACACTGGCTA CGTAGCTCCA	GCAGGIAA GACAGGIACT TCIAACTAIAA ACACTGGCIA CGIAGCICCA GATGAAA	GCAGGINA GACGGGTACB TCTAACTACAA ACACTGGCTA C	_	GGIAA GACAGGTACT TCTAACT		ACTGGYTA YGTAGCTCCA GAIG
၀၁)	# noisesen	M90528					3 8	A AB006873				9	Q.	3 9	AF159448		3 6	9	· co	la X67867		249094		9	18 x67870		1.8 A.T002290		Selected sequence for hybridization probe	colocted gemiance for	selected sequence for amplification primer
	v	o pula	atoda atoda	ptoda pho1a	atuda atuda	10 php1a		photos ploda	project plada	phola	15 obela		ploda	produ	elada elada	20 pho1a		ppo1a		800 800	25		elada elada	Blada	photo.	30 pho1a		ppja	Sele 35 hyb	2100	amp

The sequence numbering refers to the Streptococcus pneumoniae pbpla gene fragment (SEQ ID NO. 1004). Nucleotides in capitals are identical to the selected sequences or match those sequences. Mismatches are indicated by lower- case letters. Dots indicate gaps in the sequences displayed.
"-"indicates incomplete sequence data.
"R" 'Y" "W" and "S" designate nucleotide positions which are degenerated. "R" stands for A or G; "Y" stands for C or T; "W" stands for A or T; "S" stands for C or G. "I" stands for inosine which is a nucleotide analog that can bind to any of the four nucleotides A, C, G or T.
This sequence is the reverse-complement of the selected primer.

⁴⁵

Annex XXXII: Specific and ubiquitous primers for nucleic acid amplification (toxin sequences).

			Originating	DNA fragment
5	SEQ ID NO.	Nucleotide sequence	SEQ ID	Nucleotide position
0	Toxin gene:	cdtA		
	2123	5'-TCT ACC ACT GAA GCA TTA C	2129 ^a	442-460
	2124b	5'-TAG GTA CTG TAG GTT TAT TG	2129 ^a	580-599
5	Toxin gene:	cdtB	•	
	0106	5'-ATA TCA GAG ACT GAT GAG	2130 ^a	2665-2682
	2126 2127 ^b	5'-TAG CAT ATT CAG AGA ATA TTG T	2130 ^a	2746-2767
20	Toxin gene:	stx,		
	1081	5'-ATG TCA GAG GGA TAG ATC CA	1076 ^a	233-252
	1081 1080 ^b	5'-TAT AGC TAC TGT CAC CAG ACA ATG	r 1076 ^a	394-418
25	Toxin gene:	stx,		
	1078	5'-AGT TCT GCG TTT TGT CAC TGT C	1077 ^a	546-567
	1078 1079b	5'-CGG AAG CAC ATT GCT GAT T	1077 ^a	687-705
30	Toxin genes:	stx, and stx,		
	1082	5'-TTG ARC RAA ATA ATT TAT ATG TG	1076 ^a	278-300
	1082 1083 ^b	5'-TGA TGA TGR CAA TTC AGT AT	1076 ^a	781-800
35				

a Sequences from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex XXXIII: Molecular beacon internal hybridization probes for specific detection of toxin sequences.

		Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence ^a	SEQ ID NO.	Nucleotide position
Toxin gen	e: cdtA		
2125 ^b	5'-CAC GCG GAT TTT GAA TCT CTT CCT CTA GTA GCG CGT G	2129 ^c	. 462-488
Toxin ger	e: cdtB		
2128	5'- <u>CAA</u> <u>CG</u> C TGG AGA ATC TAT ATT TGT AGA AAC TGC <u>GTT</u> <u>G</u>	2130 ^C	2714-2740
Toxin ger	e: stx,		
1084	5'- <u>CCA CGC</u> CGC TTT GCT GAT TTT TCA CAT GTT ACC <u>GCG</u> TGG	1076 ^C	337-363
2012 ^d	5'- <u>CCG CGG</u> ATT ATT AAA CCG CCC TT <u>C CGC</u> <u>GG</u> -MR-HEG-ATG TCA GAG GGA TAG ATC CA	1076 ^C	248-264
Toxin ger	ne: stx,		
1085	5'- <u>CCA CGC</u> CAC TGT CTG AAA CTG CTC CTG TG CGT GG	1077 ^C	617-638
	Toxin gen 2125b Toxin gen 2128 Toxin gen 1084 2012d Toxin gen	10x111 gene: 2125b 5'-CAC GCG GAT TTT GAA TCT CTT CCT CTA GTA GCG CGT G Toxin gene: 2128 5'-CAA CGC TGG AGA ATC TAT ATT TGT AGA AAC TGC GTT G Toxin gene: 2128 5'-CCA CGC CGC TTT GCT GAT TTT TCA CAT GTT ACC GCG TGG 2012d 5'-CCG CGG ATT ATT AAA CCG CCC TTC CGC GG-MR-HEG-ATG TCA GAG GGA TAG ATC CA Toxin gene: 2012d 5'-CCG CGG CAC TGT CTG AAA CTG CTC CTG	SEQ ID NO. Nucleotide sequence Toxin gene: CdtA 2125b 5'-CAC GCG GAT TTT GAA TCT CTT CCT CTA 2129c GTA GCG CGT G Toxin gene: cdtB 2128 5'-CAA CGC TGG AGA ATC TAT ATT TGT AGA 2130c AAC TGC GTT G Toxin gene: stx, 1084 5'-CCA CGC CGC TTT GCT GAT TTT TCA CAT 1076c GTT ACC GCG TGG 2012d 5'-CCG CGG ATT ATT AAA CCG CCC TTC CGC GG-MR-HEG-ATG TCA GAG GGA TAG ATC CA Toxin gene: stx, 1085 5'-CCA CGC CAC TGT CTG AAA CTG CTC CTG 1077c

a Underlined nucleotides indicate the molecular beacon's stem.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

c Sequences from databases.

d Scorpion primer. 40

Annex XXXIV: Specific and ubiquitous primers for nucl ic acid amplification (van sequenc s).

SEQ ID NO. Nucleotide sequence SEQ ID NO. Nucleotide position				Originating DNA fragment
1086 5'-CTA CTC CCG CCT TTT GGG TT 1049-1057a 513-532b 1087c 5'-CTC ACA GCC CGA AAC AGC CT 1049-1057a 699-718b 1088c 5'-CTA CTC CCG CCT TTT GGG TT 1049-1057a 885-904b 1088c 5'-TGC CGT TTC CTG TAT CCG TC 1049-1057a 885-904b 1088c 5'-TGC CGT TTC CTG TAT CCG TC 1049-1057a 885-904b 1089c 5'-AAT AGC GGG CTAG ACC TC 1049-1057a 933-952b 1099 5'-AAT AGC GGG GAC GAA TTG GAC 1049-1057a 629-649b 1091c 5'-AAT AGC GGG GAC GAA TTG GAC 1049-1057a 629-649b 1099 5'-AAT AGC GGG GAC GAA TTG GAC 1049-1057a 629-649b 1089c 5'-ATC CAC AGG GGC TAG ACC TC 1049-1057a 629-649b 1089c 5'-ATC CAC AGG GGC TAG ACC TC 1049-1057a 629-649b 1089c 5'-ATC CAC AGG GGC TAG ACC TC 1049-1057a 629-649b 1089c 5'-ATC CAC AGG GGC TAG ACC TC 1049-1057a 662-682b 1088c 5'-TGC GGT AGC ACT TC 1049-1057a 662-682b 1088c 5'-TGC GGT AGC AGC TC 1049-1057a 885-904b 1095 5'-CGA TAG AGC GGA GAC AA 1117d 611-630 662-682b 1096c 5'-CTG ATG GAT GGG GAA GAT ACC 1049-1057a 885-904b 1096c 5'-CTG ATG GAT GGG GAA GAT ACC 1049-1057, 1117a 473-456b 1112 5'-GGC TGY GAT ATT CAA AGC TC 1049-1057, 1117a 437-456b 1114c 5'-TCW GAG CCT TTT TCC GGC TGC 1049-1057, 1117a 817-837b 1114c 5'-TCW GAG CCT TTT TCC GGC TGC 1049-1057, 1117a 705-731b 1114c 5'-TCW GAG CCT TTT TCC GGC TGG 1049-1057, 1117a 705-731b 1114c 5'-TCW GAG CCT TTT TCC GGC TGG 1049-1057, 1117a 705-731b 1114c 5'-TCW GAG CCT TTT TCC GGC TGG 1049-1057, 1117a 705-731b 1114c 5'-TCW GAG CCT TTT TCC GGC TGG 1049-1057, 1117a 817-837b 1114c 5'-TCW GAG CCT TTT TCC GGC TGG 1049-1057, 1117a 817-837b 1114c 5'-TCW GAG CCT TTT TCC GGC TGG 1049-1057, 1117a 817-837b 1114c 5'-TCW GAG CCT TTT TCC GGC TGG 1049-1057, 1117a 817-837b 1114c 5'-TCW GAG CCT TTT TCC GGC TGG 1049-1057, 1117a 817-837b 1114c 5'-TCW GAG CCT TTT TCC GGC TGG 1049-1057, 1117a 817-837b 1114c 5'-TCW GAG CCT TTT TCC GGC TGG 1049-1057, 1	5	SEQ ID NO.	Nucleotide sequence	
1087c 5'-CTC ACA GCC CGA AAC AGC CT 1049-1057a 699-718b	10	Resistanc	ce gene: vanA	
1087° 5'-CTC ACA GCC CGA AAC AGC CT 1049-1057 ^a 699-718 ^b 1086° 5'-CTA CTC CCG CCT TTT GGG TT 1049-1057 ^a 885-904 ^b 1088° 5'-TGC CGT TTC CTG TAT CCG TC 1049-1057 ^a 885-904 ^b 1089° 5'-ATC CAC ACG GGC TAG ACC TC 1049-1057 ^a 933-952 ^b 1090° 5'-AAT AGC GGG GAC GAA TTG GAC 1049-1057 ^a 734-753 ^b 1090° 5'-AAT AGC GGG GAC GAA TTG GAC 1049-1057 ^a 734-753 ^b 1090° 5'-AAT AGC GGG GAC GAA TTG GAC 1049-1057 ^a 734-753 ^b 1090° 5'-AAT AGC GGG GAC GAA TTG GAC 1049-1057 ^a 734-753 ^b 1090° 5'-AAT AGC GGG GAC GAA TTG GAC 1049-1057 ^a 933-952 ^b 1092° 5'-ATC CAC ACG GGC TAG ACC TC 1049-1057 ^a 885-904 ^b 1092° 5'-TCG GCA AGA CAA TAT GAC AGC 1049-1057 ^a 885-904 ^b 30 Resistance Gene: vanB 1095° 5'-CGA TAG AAG CAG GAC AA 1117 ^d 611-630 35 Resistance Genes: vanA, vanB 1112° 5'-GGC TGY GAT ATT CAA AGC TC 1049-1057, 1117 ^a 437-456 ^b 1112° 5'-ACC GAC CTC ACA GCC CGA AA 1049-1057, 1117 ^a 437-456 ^b 1114° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057, 1117 ^a 817-837 ^b 1116° 5'-TTC GG GCT GTG AGG TCG GBT GHG CG 1049-1057, 1117 ^a 817-837 ^b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057, 1117 ^a 705-731 ^b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057, 1117 ^a 817-837 ^b 1112° 5'-GGC TGY GAT ATT CAA AGC TC 1049-1057, 1117 ^a 817-837 ^b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057, 1117 ^a 705-731 ^b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057, 1117 ^a 817-837 ^b 1112° 5'-GGC TGY GAT ATT CAA AGC TC 1049-1057, 1117 ^a 817-837 ^b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057, 1117 ^a 817-837 ^b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057, 1117 ^a 817-837 ^b 1112° 5'-GGC TGY GAT ATT CAA AGC TC 1049-1057, 1117 ^a 817-837 ^b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057, 1117 ^a 817-837 ^b 1112° 5'-GGC TGY GAT ATT CAA AGC TC 1049-1057, 1117 ^a 817-837 ^b 11112° 5'-GGC TGY GAT ATT CAA AGC TC 1049-1057, 1117 ^a 817-837 ^b		1006	SY_CTA CTC CCG CCT TTT GGG TT	1049-1057 ^a 513-532 ^b
1086		_ '		1049-1057 ^a 699-718 ^b
1086			and and account man account	1049-1057 ^a 513-532 ^b
1086 5'-GC CGT TTC CGG CCT TTT GGG TT 1049-10578 513-532b 1089° 5'-ATC CAC ACG GGC TAG ACC TC 1049-10579 933-952b 1091° 5'-AAC GCG GAC GAA TTG GAC 1049-10579 629-649b 1091° 5'-AAC GCG GCA CTG TTT CCC AA 1049-10579 734-753b 1089° 5'-ATC CAC ACG GGC TAG ACC TC 1049-10579 734-753b 1089° 5'-ATC CAC ACG GGC TAG ACC TC 1049-10579 933-952b 1089° 5'-ATC CAC ACG GGC TAG ACC TC 1049-10579 933-952b 1089° 5'-TCG GCA AGA CAA TAT GAC AGC 1049-10579 933-952b 1088° 5'-TCG CGT TTC CTG TAT CCG TC 1049-10579 885-904b 1088° 5'-TCG CGT TTC CTG TAT CCG TC 1049-10579 885-904b 1096° 5'-CTG ATG GAT GCG GAC AA 1117d 611-630 885-904b 1112° 5'-GCG TGY GAT ATT CAA AGC TC 1049-1057,11179 437-456b 1113° 5'-ACC GAC CTC ACA GCC CGA AA 1049-1057,11179 705-724b 1114° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,11179 817-837b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,11179 817-837b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,11179 817-837b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,11179 817-837b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,11179 817-837b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,11179 817-837b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,11179 817-837b 1112° 5'-GCC TGY GAT ATT CAA AGC TC 1049-1057,11179 817-837b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,11179 817-837b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,11179 817-837b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,11179 817-837b 1112° 5'-GCC TCY GAT ATT CAA AGC TC 1049-1057,11179 817-837b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,11179 817-837b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,11179 817-837b 1116° 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,11179 817-837b 1112° 5'-GCC TCY GAT ATT CAA AGC TC 1049-1057,11179 817-837b 1112° 5'-GCC TCY GAT ATT CAA AGC TC 1049-1057,11179 817-837b 1112° 5'-GCC TCY GAT ATT CAA AGC TC 1049-1057,11179 817-837b	15			1
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1114 ^C 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,1117 ^a 817-837 ^b 1116 5'-TTT CGG GCT GTG AGG TCG GBT GHG CGG 1049-1057,1117 ^a 705-731 ^b 1114 ^C 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,1117 ^a 817-837 ^b 1112 5'-GGC TGY GAT ATT CAA AGC TC 1049-1057,1117 ^a 437-456 ^b		1114-	3 -16W GAG CC1 111 100 000 100	
1116 5'-TTT CGG GCT GTG AGG TCG GBT GHG CGG 1049-1057,1117a 705-731b 1114c 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,1117a 817-837b 1112 5'-GGC TGY GAT ATT CAA AGC TC 1049-1057,1117a 437-456b		1115		1049-1057,1117ª 705-730 ^D
1116 5'-TTT CGG GCT GTG AGG TCG GBT GHG CGG 1049-1057,1117a 705-731b 1114c 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,1117a 817-837b 1112 5'-GGC TGY GAT ATT CAA AGC TC 1049-1057,1117a 437-456b		1114 ^C	5'-TCW GAG CCT TTT TCC GGC TCG	1049-1057,1117ª 817-837 [©]
1114 ^C 5'-TCW GAG CCT TTT TCC GGC TCG 1049-1057,1117 ^a 817-837 ^b 1112 5'-GGC TGY GAT ATT CAA AGC TC 1049-1057,1117 ^a 437-456 ^b	:5	1116	EVENTE COG OCT OTG AGG TOG GBT GHG CGG	1049-1057,1117 ^a 705-731 ^b
1112 5'-GGC TGY GAT ATT CAA AGC TC 1049-1057,1117 ^a 437-456 ^b			5'-TCW GAG CCT TTT TCC GGC TCG	1049-1057,1117 ^a 817-837 ^b
1112 3 000 101 0112 1112		****		1040 1057 11178 437-456b
1118° 5'-TTT TCW GAG CCT TTT TCC GGC TCG 1043-103/,111. 01. 01.			5'-GGC TGY GAT ATT CAA AGC TC	1049-1057,1117ª 817-840b
	:0	1118 ^C	5'-TTT TCW GAG CCT TTT TCC GGC TCG	1045 1057,1117

These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the vanA sequence fragment (SEQ ID NO. 1051).

These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

d Sequences from databases.

Specific and ubiquitous primers for nucleic acid Annex XXXIV: amplification (van s quences) (continued).

			Originating DNA fragment
5	SEQ ID NO.	Nucleotide sequence	SEQ ID Nucleotide NO. position
10	Resistan	ce genes: vanA, vanB (continu	ied)
	1115	5'-TTT CGG GCT GTG AGG TCG GBT GHG CG	1049-1057,1117 ^a 705-730 ^b
	1118 ^C	5'-TTT TCW GAG CCT TTT TCC GGC TCG	1049-1057,1117 ^a 817-840 ^b
1.5	1116	5'-TTT CGG GCT GTG AGG TCG GBT GHG CGG	1049-1057,1117 ^a 705-731 ^b
15	1118 ^C	5'-TTT TCW GAG CCT TTT TCC GGC TCG	1049-1057,1117 ^a 817-840 ^b
			1049-1057,1117 ^a 705-729 ^b
	1119	5'-TTT CGG GCT GTG AGG TCG GBT GHG C 5'-TTT TCW GAG CCT TTT TCC GGC TCG	1049-1057,1117 ^a 817-840 ^b
20	1118 ^C		
	1120	5'-TTT CGG GCT GTG AGG TCG GBT GHG	1049-1057,1117ª 705-728b
	1118 ^C	5'-TTT TCW GAG CCT TTT TCC GGC TCG	1049-1057,1117 ^a 817-840 ^b
	1121	5'-TGT TTG WAT TGT CYG GYA TCC C	1049-1057,1117 ^a 408-429 ^b
25	1111 ^C	5'-CTT TTT CCG GCT CGW YTT CCT GAT G	1049-1057,1117 ^a 806-830 ^b
		THE TOTAL COMPANY CAN ACCURE	1049-1057,1117 ^a 437-456 ^b
	1112	5'-GGC TGY GAT ATT CAA AGC TC 5'-CTT TTT CCG GCT CGW YTT CCT GAT G	1049-1057,1117 ^a 806-830 ^b
	1111 ^c		
30	1123	5'-TTT CGG GCT GTG AGG TCG GBT G	1049-1057,1117 ^a 705-726 ^b 1049-1057,1117 ^a 806-830 ^b
	1111 ^c	5'-CTT TTT CCG GCT CGW YTT CCT GAT G	1049-1057,11174 806-8304
	1112	5'-GGC TGY GAT ATT CAA AGC TC	1049-1057,1117 ^a 437-456 ^b
	1124 ^C	5'-GAT TTG RTC CAC YTC GCC RAC A	1049-1057,1117 ^a 757-778 ^b
35	Resistar	nce gene: vanCl	
	1103	5'-ATC CCG CTA TGA AAA CGA TC	1058-1059 ^a 519-538 ^d
	1103 1104 ^C	5'-GGA TCA ACA CAG TAG AAC CG	1058-1059 ^a 678-697 ^d
40		nce genes: vanCl, vanC2, vanC	3
	1007	5'-TCY TCA AAA GGG ATC ACW AAA GTM AC	1058-1066 ^a 607-632 ^d
	1097	5'-TCT TCA AAA TCG AAA AAG CCG TC	1058-1066 ^a 787-809 ^d
45	1098 ^C	57-TCT TCA AAA TCG AAA AAG CCG TC	
	1099	5'-TCA AAA GGG ATC ACW AAA GTM AC	1058-1066 ^a 610-632 ^d
	1100 ^C	5'-GTA AAK CCC GGC ATR GTR TTG ATT TC	1058-1066 ^a 976-1001 ^d
	1101	5'-GAC GGY TTT TTY GAT TTT GAA GA	1058-1066 ^a 787-809 ^d
50	1101 ^c	5'-AAA AAR TCG ATK CGA GCM AGA CC	1058-1066 ^a 922-944 ^d
• •	Resistanc		
	1105	5'-CTC CTA CGA TTC TCT TGA YAA ATC A	1060-1066,1140 ^a 487-511 ^e
55	1105 1106 ^C	5'-CAA CCG ATC TCA ACA CCG GCA AT	1060-1066,1140 ^a 690-712 ^e
,,,			

a These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the vanA sequence fragment (SEQ ID NO. 1051).

These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

d The nucleotide positions refer to the vanCl sequence fragment (SEQ ID NO. 1058).

e The nucleotide positions refer to the vanC2 sequence fragment (SEQ ID NO. 1140).

Annex XXXIV: Specific and ubiquitous primers for nucleic acid amplification (van sequences) (continued).

			Originating DNA fragmen
5	SEQ ID NO.	Nucleotide sequence	SEQ ID Nucleotide NO. position
10	Resistanc	ce gene: vanD	
	1591	5'-ATG AGG TAA TAG AAC GGA TT	1594 797-837
	1592 ^b	5'-CAG TAT TTC AGT AAG CGT AAA	1594 979-999
5	Resistanc	ce gene: vanE	
	1595	5'-AAA TAA TGC TCC ATC AAT TTG CTG A	1599 ^a 74-98
	1596 ^b	5'-ATA GTC GAA AAA GCC ATC CAC AAG	1599 ^a 394-417
0	1597	5'-GAT GAA TTT GCG AAA ATA CAT GGA	1599 ^a 163-186
U	1598 ^b	5'-CAG CCA ATT TCT ACC CCT TTC AC	1599 ^a 319-341
		Sequencing primers	(vanAB)
25	1112	5'-GGC TGY GAT ATT CAA AGC TC	1139 ^a 737-756
,	1111 ^b	5'-CTT TTT CCG GCT CGW YTT CCT GAT G	1139 ^a 1106-1130
		Sequencing primers	(vanA, vanX, vanY)
80	1150	5'-TGA TAA TCA CAC CGC ATA CG	1141 ^a 860-879
U	1151 ^b	5'-TGC TGT CAT ATT GTC TTG CC	1141 ^a 1549-1568
	1152	5'-ATA AAG ATG ATA GGC CGG TG	1141 ^a 1422-1441
	1152 1153b	5'-CTC GTA TGT CCC TAC AAT GC	1141a 2114-2133
35	1154	5'-GTT TGA AGC ATA TAG CCT CG	1141 ^a 2520-2539
,,,	116/		

a Sequences from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex XXXIV: Sp cific and ubiquitous primers for nucleic acid amplification (van sequences) (continued).

			Originating	DNA fragment
5	SEQ ID NO.	Nucleotide sequence SEQ ID	Nucleotide NO.	position
10		Sequencing primers	(vanC1)	
	1110 1109 ^b	5'-ACG AGA AAG ACA ACA GGA AGA CC 5'-ACA TCG TGA TCG CTA AAA GGA GC	1138 ^a 1138 ^a	122-144 1315-1337
15		Sequencing primers	(vanC2, vanC	3)
	1108 1107 ^b	5'-GTA AGA ATC GGA AAA GCG GAA GG 5'-CTC ATT TGA CTT CCT CCT TTG CT	1140 ^a 1140 ^a	1-23 1064-1086
20				

a Sequences from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex XXXV: Internal hybridization probes for sp cific detection of van sequences.

		Originating	DNA fragment
SEQ ID 1	NO. Nucleotide sequence	SEQ ID NO.	Nucleotide position
Resist	ance gene: vanA		
1170 2292	5'-ACG AAT TGG ACT ACG CAA TT 5'-GAA TCG GCA AGA CAA TAT G	1049-1057 ^a 2293 ^c	639-658 ^b 583-601
Resist	ance gene: vanB		
1171 2294 2295	5'-ACG AGG ATG ATT TGA TTG TC 5'-AAA CGA GGA TGA TTT GAT TG 5'-TTG AGC AAG CGA TTT CGG	1117 ^C 2296 ^a 2296 ^a	560-579 660-679 614-631
Resist	ance gene: vanD		
2297	5'-TTC AGG AGG GGG ATC GC	1594 ^C	458-474

a These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the vanA sequence fragment (SEQ ID NO. 1051).

c Sequences from databases.

Annex XXXVI: Specific and ubiquitous primers for nucleic acid amplification (pbp sequences).

			Originating	DNA fragment
5	SEQ ID NO.	Nucleotide sequence	SEQ ID. NO.	Nucleotide position
10	Resistance	gene: pbpla		
	1129	5'-ATG ATG ACH GAM ATG ATG AAA AC	1004-1018 ^a	681-703 ^b
	1131 ^C	5'-CAT CTG GAG CTA CRT ARC CAG T	1004-1018 ^a	816-837 ^b
15	1130	5'-GAC TAT CCA AGC ATG CAT TAT G	1004-1018 ^a	456-477 ^b
15	1131	5'-CAT CTG GAG CTA CRT ARC CAG T	1004-1018 ^a	816-837 ^b
	2015	5'-CCA AGA AGC TCA AAA ACA TCT G	2047 ^d	909-930
	2015 2016 ^C	5'-TAD CCT GTC CAW ACA GCC AT	2047 ^d	1777-1796
20		Sequencing primers	(pbpla)	
		-		072 002
	1125	5'-ACT CAC AAC TGG GAT GGA TG	1169 ^d 1169 ^d	873-892 2140-2160
25	1126 ^C	5'-TTA TGG TTG TGC TGG TTG AGG	1163-	2140-2100
23	1125	5'-ACT CAC AAC TGG GAT GGA TG	1169 ^d	873-892
	1128 ^C	5'-GAC GAC YTT ATK GAT ATA CA	1169 ^d	1499-1518
	1127	5'-KCA AAY GCC ATT TCA AGT AA	1169 ^d	1384-1403
30	1126 ^C	5'-TTA TGG TTG TGC TGG TTG AGG	1169 ^d	2140-2160
		Sequencing primers	(pbp2b)	
		5'-GAT CCT CTA AAT GAT TCT CAG GTG		1-25
35	1142 1143 ^C	5'-CAA TTA GCT TAG CAA TAG GTG TTG	-	1481-1505
22	1143			4 05
	1142	5'-GAT CCT CTA AAT GAT TCT CAG GTG	G 1172 ^d	1-25
	1145 ^C	5'-AAC ATA TTK GGT TGA TAG GT	1172 ^d	793-812
40	1144	5'-TGT YTT CCA AGG TTC AGC TC	1172 ^d	657-676
	1143 ^C	5'-CAA TTA GCT TAG CAA TAG GTG TTG	G 1172 ^d	1481-1505
		Sequencing primers (pbp2x)	
15	1146	5'-GGG ATT ACC TAT GCC AAT ATG AT	1173 ^d	219-241
	1143 1147 ^C	5'-AGC TGT GTT AGC VCG AAC ATC TTG	1173 ^d	1938-1961
		TA GOO NEW ACC MAN COC ARE ACC AC	1173 ^d	219-241
50	1146 1149 ^C	5'-GGG ATT ACC TAT GCC AAT ATG AT 5'-TCC YAC WAT TTC TTT TTG WG	1173d	1231-1250
,0	****		11m2d	711 720
	1148_	5'-GAC TTT GTT TGG CGT GAT AT	1173 ^d 1173 ^d	711-730 1938-1961
	1147 ^C	5'-AGC TGT GTT AGC VCG AAC ATC TTG	11/3~	1330-1301

^a These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the pbpla sequence fragment (SEQ ID NO. 1004).

C These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

j0 d Sequences from databases.

Annex XXXVII: Internal hybridization probes for specific detection of pbp s quences.

5		Origina	ting DNA fragment
	SEQ ID NO.	Nucleotide sequence SEQ 1	D Nucleotide position
10	Resistance	gene: pbpla	
	1132	5'-AGT GAA AAR ATG GCT GCT GC 1004-10	
	1132	5'-CAT CAA GAA CAC TGG CTA YGT AG 1004-10	
15	1134	5'-CTA GAT AGA GCT AAA ACC TTC CT 1004-10	018 ^a 417-439 ^b
13	1135	5'-CAT TAT GCA AAC GCC ATT TCA AG 1004-10	
	1192	5'-GGT AAA ACA GGA ACC TCT AAC T 1004-10	
	1193	5'-GGT AAG ACA GGT ACT TCT AAC T 1004-1	
	1194	5'-CAT TTC AAG TAA TAC AAC AGA ATC 1004-1	018 ^a 485-508 ^b
20	1195	5'-CAT TTC AAG TAA CAC AAC TGA ATC 1004-1	018 ^a 485-508 ^b
20	1196	5'-GCC ATT TCA AGT AAT ACA ACA GAA 1004-1	018 ^a 483-506 ^b
	1197	5'-CAA ACG CCA TTT CAA GTA ATA CAA C 1004-1	018 ^a 478-502 ^b
	1094	5'-GGT AAA ACA GGT ACT TCT AAC TA 1004-1	018 ^a 759-781 ^b
	1214	5'-GGT AAA ACA GGT ACC TCT AAC TA 1004-1	018 ^a 759-781 ^b
25	1214	5'-GGT AAG ACT GGT ACA TCA AAC TA 1004-1	018 ^a 759-781 ^b
23	1217	5'-CAA ATG CCA TTT CAA GTA ACA CAA C 1004-1	
	1218	5'-CAA ACG CCA TTT CAA GTA ACA CAA C 1004-1	018 ^a 478-502 ^b
	1219	5'-CAA ATG CTA TTT CAA GTA ATA CAA C 1004-1	018 ^a 478-502 ^b
	1220	5'-CAA ACG CCA TTT CAA GTA ATA CGA C 1004-1	
30	2017	5'-ACT TTG AAT AAG GTC GGT CTA G 204'	7 ^C 1306-1327
50	2017	5'-ACA CTA AAC AAG GTT GGT TTA G 206	3 354-375
	2019	5'-ACA CTA AAC AAG GTC GGT CTA G 206	4 346-367
	2020	5'-GTA GCT CCA GAT GAA ATG TTT G 214	oc 1732-1753
	2021	5'-GTA GCT CCA GAC GAA ATG TTT G 205	831-852
35	2022	5'-GTA GCT CCA GAT GAA ACG TTT G 205	3 ^c 805-826
55	2023	5'-GTA ACT CCA GAT GAA ATG TTT G 205	819-840
	2024	5'-AGT GAA AAG ATG GCT GCT GC 204	8 ^C 1438-1457
	2025	5'-AGT GAG AAA ATG GCT GCT GC 204	7 ^C 1438-1457
	2026	5'-TCC AAG CAT GCA TTA TGC AAA CG 204	7 ^C 1368-1390
•		5'-TCG GTC TAG ATA GAG CTA AAA CG 2047C	1319-1341
0	2027 2028	5'-TAT GCT CTT CAA CAA TCA CG 2047C	
	2028	5'-AGC CGT TGA GAC TTT GAA TAA G 2047C	
	2029	5'-CTT AAT GGT CTT GGT ATC G 2047C	
	2030	5'-CGT GAC TGG GGT TCT GCT ATG A 2049C	
15	2031	5'-CGT GAC TGG GGA TCA TCA ATG A 2047C	1096-1117
•3	2032	5'-CGT GAC TGG GGT TCT GCC ATG A 2057	195-216
	2033	5'-ATC AAG AAC ACT GGC TAT GTA G 2050°	787-808
	2034		

⁵⁰ a These sequences were aligned to derive the corresponding primer.

b The nucleotide positions refer to the pbpla sequence fragment (SEQ ID NO. 1004).

c Sequence from databases.

Annex XXXVII: Internal hybridization probes for specific detection of pbp sequences (continued).

				Originatin	g DNA fragmen
SEQ ID NO.	Nucleotid	e sequenc	e	SEQ ID NO.	Nucleotide position
Resistance	gene:	pbp1a	(continued)		
2035	5'-ATC AA	G AAC ACT	GGC TAC GTA G	2051 ^C	787-808
2036	5'-ATC AA	G AAC ACT	GGT TAC GTA G	2047	1714-1735
2037	5'-ATC AA	A AAT ACT	GGT TAT GTA G	2057	813-834
2038			GGC TAC GTA G	2052 ^C	757-778
2039			GGC TAT GTA G	2053 ^C	787-808

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SEQ ID NO.: 1139 1141 1051 1053	1054 1055 1055 1056 1057	1050 1117 -		1 1 1 1 1	1112
NGCCGGACGAATT GGACTACGCA NGCCGGACGAATT GGACTACGCA NGCCGGACGAATT GGACTACGCA NGCCGGACGAATT GGACTACGCA	GCGATATICA AAGCTCAGCCGGACGAATT GCGATATICA AAGCTCAGCCGGACGAATT GCGATATICA AAGCTCAGCCGGACGAATT GCGATATICA AAGCTCAGCCGGACGAATT GCGATATICA AAGCTCAGCCGGACGAATT	GIAGGCT GCGATATY CA AAGCTCAGCCGGACGAAIT GGACTACGCA AILGAA GIGGGCT GTGATATY CA AAGCTCAGCCGGACGAAIT GGACTACGCA ATTGAA GIAGGCT GCGATATY CA AAGCTCCGCCGGAAGAACT tAACGCTGCG ATGGAA GIGGGCT GTGATATY CA AAGCTCCGCCGGAAGAACT BAACGCTGCG ATAGAA	GTGATAT! CA AAGCTCCGCCGGAAGAACT GTGATAT! CA AAGCTCCGCCGGAAGAACT GTGATAT! CA AAGCTCCGCCGGAAGAACT GCGATAT! CA AAGCTCCGCCGGAAGAACT GCGATAT! CA AAGCTCCGCCGGAAGAACT	GTAGGCT GCGATATICA AAGCTCCGCCGGAAGAACT taANGGCTGU AIRGAA GTGGGCT GTGATATICA AAGCTCCGCCGGAAGAACT BAANGGCGG ATBGAA GTAGGCT GCGATATICA AAGCTCCGCCGGAAGAACT taANGGCGG ATBGAA GTGGGAT GCGATATICA AAGCTCCGCCGGAAGAACT BAANGGCGG ATBGAA GTAGGLT GTGGTATICA AAGCTCCGTCAGAAGAACT GCAGGCAGCA ATGGAA	GGCT GYGATAT! ICA AAGCTC ACGAATT GGACTACGCA ATT (VADA)
Accession # X56895 M97297			U94539 U94530 283305 U81452 U35369	L06138 L15304 U00456 AF130997 AF136925	Selected sequence for amplification primer Selected sequence for hybridization probe
5 vanA vanA vanA vanA	10 vana vana vana vana	15 vana vana vana	vans 20 vans vans vans vans vans	25 vanB vanB vanB vanB vanD vanB	

"R" "Y" "M" "K" "W" and "S" designate nuk stands for A or C; "K" stands for G or T;:leotide positions which are degenerated. "R" stands for A or G; "Y" stands for C or T; "M" analog that can bind to any of the four nu "W" stands for A or T; "S" stands for C or G. "I" stands for inosine which is a nucleotide cleotides A, C, G or T.

The sequence numbering refers to the Ente to the Ente faecium vanA gene fragment (SEQ ID NO. 1139). Nucleotides in capitals are identical to the selected sequences or match those srococcus faecium vanA gene fragment (SEQ ID NO. 1139). Nucleotides in capitals are identicated by lower-case letters. Dots indicate gaps in the sequences displayed.

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the selection of vanAB-specific amplification primers and sednences from van specific hybridization probes ranB-Strategy for (continued). vanA- and Annex XXXVIII:

SEQ ID NO: 1139 1141 1051 1052 1053 1056 1056 1057 1050 1117	1171	1111
GAAACAG GCGGGTT ag TTGTTGGCATT CATCAGGAAG TCGAGCCGGA AAAAGGCT GAAACAG GCGGGTT ag TTGTTGGCATT CATCAGGAAG TCGACCCGGA AAAAGGCT GAAACAG GATGATTT ag TTGTTCGCCATT CATCAGGAAG TCGACCCGGA AAAAGGCT GAAACAG GATGATTT ag TTGTTCGCCATT CATCAGGAAA ACGACCGGA AAAAGGCT GGAACGAG GATGATTT GA TTGTCGGCATT CATCAGGAAA ACGACCCGGA AAAAGGCT GGAACGAG GATGATTT GA TTGTCGGCATT CATCAGGAAA ACGACCCGGA AAAAGGCT GGAACGAG GATGATTT GA TTGTCGGCATC CATCAGGAAA ACGACCCGGA AAAAGGCT GGAACGAG GATGATTT GA TTGTCGGCATC CATCAGGAAA ACGAGCCGGA AAAAGGCT GGAACCAG GATGATTT GA TTGTCGGCATC CATCAGGAAA ACGAGCCGGA AAAAGGCT GAAACGAG GATGATTT GA TTGTCGGCATC CATCAGGAAA ACGAGCCGGA AAAAGGCT GAAACGAG GATGATTT GA TTGTCGGCATC CATCAGGAAA ACGAGCCGGA AAAAGGCT GAAACGAG GATGATTT GA TTGTCGCCATC CATCAGGAAA ACGAGCCGGA AAAAGGCT GAAACGAG GATGATTT GA TTGTCGCCATC CATCAGGAACAAACAAA ACGAGCCGGA AAAACGAAACAAAAAAAA	ACGAG GATGATTI GA TTGTC (VALB)	CATCAGGAAR WCGAGCCGGA AAAAG
vanA X56895 vanA X56895 vanA M97297 vanA vanA vanA vanA vanB U94526 vanB U94527 vanB U94529 vanB U94529 vanB U94529 vanB U94529 vanB U94529 vanB U04529 vanB U01704 vanB L15304 vanB L15304 vanB U00456 vanB AF136925	Selected sequence for hybridization probe	Selected sequence for amplification primer
S 0 51 8 8 0E		35

The sequence numbering refers to the Enter faecium vanA gene fragment (SEQ ID NO. 1139). Nucleotides in capitals are identical to the selected sequences or match those se ococcus faecium vanA gene fragment (SEQ ID NO. 1139). Nucleotides in capitals are identical displayed. which are degenerated. "R" stands for A or G; "W" stands for A or T displayed. "R" and "W" designate nucleotide positions

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of the above selected primer. This sequence is the reverse-complement

	Annex XXXIX:	Int rnal detection	hybridization of mecA.	probe	for	specific
				Origi	nating	DNA fragment
5	SEQ ID NO.	Nucleotide se	equence		D ID	Nucleotide position
10	Resistance g	ene:	necA			
	1177	5'-GCT CAA C	AA GTT CCA GAT TA	11	78 ^a	1313-1332

a Sequence from databases.

Specific and ubiquitous primers for nucleic acid Annex XL: amplification (hexA sequences).

5								Originating	DNA fragment
	SEQ ID NO		Nucleon	ide s	equenc	e		SEQ ID NO.	Nucleotide position
10	Bacteria	l spec	cies:		Strep	tococ	cus pneumo	nise	
		5'-ATT 5'-AGC						1183 ^a 1183-1191 ^c	431-450 652-671 ^d
15					Seque	ncing	primers		•
20		5'-ATT 5'-AAC						1183 ^a 1183 ^a	431-450 1045-1064

a Sequences from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing. 25

C These sequences were aligned to derive the corresponding primer.

d The nucleotide positions refer to the hexA sequence fragment (SEQ ID NO. 1183).

Annex XLI: Internal hybridization probe for specific detection of hexA sequences.

5			Originating	DNA fragment
	SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
10	Bacterial s	species: Streptococcus pn	eumoniae	
	1180 ^a	5'-TCC ACC GTT GCC AAT CGC A	1183-1191 ^b	629-647 ^C
15				

a This sequences is from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

b These sequences were aligned to derive the corresponding primer.

²⁰ C The nucleotide positions refer to the hexA sequence fragment (SEQ ID NO. 1183).

amplification prie selection of Streptococcus pneumoniae species-specific imers and hybridization probe from hexA sequences. Ann x XLII: Strategy for

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		-		SEQ ID
	*	428	1067	NO.:
	S. pneumoniae	TGG ATTTOGTONC GOGTONCTT	453 626 CONTRACTOR DESCRIPTION AND ADDRESS AND ADDRESS	1183
01		TOAC GGGTGACTTT'	TATATTTG CGATTGGCAA CGGTGGAGCA AACGGCATCT AGTAAGCTGC TCCAAATCTABAAG GATCTCTTG	1184
		TOAC GGOTOACTTT	TATATTTG CGATTGGCAA CCGTGGAGCA AACGGCAICT AGTAAGCISC LCCAAATTG	1185
٠		TOAC GGOTGACTTT	TATATTTG CGATTGCCAA CCGTGGGGCA AACGGCALCI AGIAAACICC CCCCAITTG CGATTGCCAA AACGCAACIC AACGCAACAA AACGCAAGA AACGAAGA AACGCAAGA AACGCAAGA AACGCAAGA AACGCAAGA AACGCAAGA AACGCAAGA AACGCAAGA AACGCAAGA AACCAAGA AACCAAGAAACAACAAACA	1186
		TGAC GGGTGACTTT	TAT ATTTG CGATTGGCAA CGGTGGGGCA AACGGCATUT AGTAAGTGC TUCA: AATGCAAAG GATCTCTT	1187
	S. oralis	GGGTGACTTT	TAT ATTITO COATTGGGAA CGGIGGAGGA MACGGGALCA SCANGAGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	1188
15		GOTGAC GGGTGACTTT	TAT ATCCA CGACTGGCAG CCUTGGAGLA AGCGGCAGCA AGAMACACACA AGACCAAAG GATCTCTT	1189
2		TOAC OGGTGACTTI	TAT ATTER COATTGGCAG CTGTGGACCA AGCGGCATCT ACTARACTEC ALCA	1190
116		TOAC GOGTGACTTT	CAGGCGag gagerdtete etarggagee Teaggedgea gegaaatie teen.	1191
			CAGGCGAG gaAcTGtCtc CtaTGGAGCG TCaGGCAGCG gGGAAAttGC thon	
20	Selected sequence for amplification primer	ATTTGGTGAC GGGTGACTTT		1179
	Selected sequences for amplification primers*			Ċ
25			ACGGCATCT AGTAAGCTGC T CCAAAG GATCTCTTGC AGTT	1181
	Selected sequence for hybridization probe*		THE CREW CONTON	1180
30				

The sequence numbering refers to the *Streptoco* sequences. ^Mccus pneumoniae hexa gene fragment (SEQ ID NO. 1183). Nucleotides in capitals are identical to the sequences or match those sequences. ^Mccus pneumoniae hexa gene fragment (SEQ ID NO. 1183). Nucleotides in the sequences data.

35

s selected primer.

This sequence is the reverse-complement of the

Annex XLIII: Specific and ubiquitous primers for nucleic acid amplification (pcp sequence).

		Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
Bacterial s	species: Streptococcus pyogen	es	
1211 1210 ^b	5'-ATT CTT GTA ACA GGC TTT GAT CCC 5'-ACC AGC TTG CCC AAT ACA AAG G	1215 ^a 1215 ^a	291-314 473-494

a Sequences from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Sp cific and ubiquitous primers for nucleic acid Annex XLIV: amplification of S. saprophyticus sequences of unknown coding potential.

		Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
Bacterial s	pecies: Staphylococcus sapro	phyticus	
1208	5'-TCA AAA AGT TTT CTA AAA AAT TTA	74,1093, 1198 ^b	169-193 ^C
1209 ^a	5'-ACG GGC GTC CAC AAA ATC AAT AGG	A 74,1093. 1198 ^b	355-379 ^C

a This sequence is from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

b These sequences were aligned to derive the corresponding primer.

 $^{^{} extsf{C}}$ The nucleotide positions refer to the S. saprophyticus unknown gene sequence fragment (SEQ ID NO. 1198).

Molecular beacon internal hybridization probes for specific d tection of antimicrobial ag nts Annex XLV: resistance gene sequences.

		riginating	DNA fragment
SEQ ID NO.	Nucleotide sequence ^a	SEQ ID	Nucleotide position
Resistanc	ce gene: gyrk		
2250	5'-CCG TCG GAT GGT GTC GTA TAC CGC GGA GTC GCC GAC GG	1954 ^b	218-243
2251	5'-CGG AGC CGT TCT CGC TGC GTT ACA TGC TGG TGG CTC CG	1954 ^b	259-286
Resistanc	ce gene: mecA		
1231	5'-GCG AGC CCG AAG ATA AAA AAG AAC CTC TGC TGC TCG C	1178 ^b	1291-1315
Resistan	ce gene: parC		
1938 ^b	5'-CCG CGC ACC ATT GCT TCG TAC ACT GAG GAG TCT CCG CGC GG	1321 ^C	232-260
1939	5'- <u>CGA CCC GG</u> A TGG TAG TAT CGA TAA TGA TCC GCC AGC GG <u>C</u> <u>CGG</u> <u>GTC</u> <u>G</u>	1321 ^c	317-346
1955 ^b	5'- <u>CGC GCA</u> ACC ATT GCT TCG TAC ACT GAG GAG TC <u>T</u> <u>GCG</u> <u>C</u> G	1321 ^c	235-260
Resistan	ice gene: vanA		
1239	5'- <u>GCG</u> <u>AGC</u> GCA GAC CTT TCA GCA GAG GAG <u>GCT</u> <u>CGC</u>	1051	860-880
1240	5'- <u>GCG AGC</u> CGG CAA GAC AAT ATG ACA GCA AAA TC <u>G CTC GC</u>	1051	663-688
Daeietan Ne <u>sistant</u>	TOO TONG! TEMP	•	
1241	5'-GCG AGC GGG GAA CGA GGA TGA TTT GAT TGG CTC GC	1117	555-577
Resistanc	<u>e gene</u> : van D		
1593	5'-CCG AGC GAT TTA CCG GAT ACT TGG CTG ICG CTC GG	1594	835-845

a Underlined nucleotides indicate the molecular beacon's stem.

b This sequence is from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

c Sequence from databases.

Annex XLVI: Molecular beacon internal hybridization probe for specific detection of S. aureus gene sequences of unknown coding pot ntial.

		Originating	Originating DNA fragment		
SEQ ID NO.	Nucleotide sequence ^a	SEQ ID	Nucleotide position		
Bacterial	species: S. aureus				
1232	5'-GGA GCC GCG CGA TTT TAT AAA TGA ATG ATA ACC GGC TCC	TTG 1244	53-80		

a Underlined nucleotides indicate the molecular beacon's stem.

Annex XLVII: Molecular beacon internal hybridization probes for specific det ction of tuf sequences.

				Originating	DNA fragment
SEQ ID NO.	Nucleotide	sequence ^a		SEQ ID NO.	Nucleotide position
Bacterial	species:	Chlamy	dia pneumoniae	•	
2091	5'-CGC GAC TTC TTG	TTG AGA TGG A	AC TTA GTG AGC	20	157-183
2092	5'-CGC GAC TGC AGG	GAA AGA ACT T	CC TGA AGG TCG	20	491-516
<u>Bacterial</u>	species:	Chlamy	dia trachomati	.	
2213	5'- <u>CGT</u> <u>GCC</u> GAC GCT	ATT GAC ATG A	TT TCC GAA GAA	1739 ^b	412-441
Bacterial	species:	Entero	coccus faecali	s	
1236	5'-GCG AGC GGC TCG		GTT CGC GTT GGT	883	370-391
<u>Bacterial</u>	species:	Entero	coccus faecium		
1235	5'-GCG AGC TGC TGG	CGA AGT TGA A	AGT TGT TGG TAT	64	412-437
<u>Bacteria</u>	<u>species</u> :	Legion	nella pneumophi	la	
2084 ^C	5'- <u>CAC GCC</u> TTT TGC	TCA ACA CCC (GTA CAA GTC GTC	112	461-486
Bacteria	l species:	Мусор	lasma pneumonia	e	
2096 ^C	5'- <u>CGC GAC</u> T <u>GT CGC</u>	CGG TAC CAC	GGC CAG TAA TCG	2097 ^b	658-679
<u>Bacteria</u>	l species:	Neiss	eria gonorrhoes	ie	
2177	5'- <u>GGC ACG</u> ATC GAA	GAC AAA CCA TT ACG TGT TC <u>C</u>	C CTG CTG CCT	126	323-357
2178	5'-GGC ACG TCG AAC	ACA AAC CAT TO	C TGC TGC CTA	126	323-348
2179	5'- <u>GGC AGC</u> TAA CCG	TCT ACT TCC GT	FA CCA CTG ACG	126	692-718

a Underlined nucleotides indicate the molecular beacon's stem.

b Sequence from databases.

C This sequence is from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex XLVII: Molecular b acon internal hybridization probes for specific d t ction of tuf sequences (continued).

			Originating	DNA fragment			
SEQ ID NO.	Nucleotide sequence ^a		SEQ ID NO.	Nucleotide position			
Bacterial	species: Pseudomona						
2122	5'-CCG AGC GAA TGT AGG AGT CC CTG CTC GG	A GGG TCT	153,880,2138	b,c 280-302 ^d			
Bacterial	species: Staphyloco	ccus aureus	•				
2186				615-646			
Bacterial	group: Staphyloco	ccus sp. ot	her than S	. aureus			
1233	5'-GCG AGC GTT ACT GGT GTA G CGG CTC GC						
Fungal_sp	ecies: Candida al	bicans					
2073			408	404-429			
Fungal sp	Fungal species: Candida dubliniensis						
2074	5'-CCG AGC AAC ATG ATT GAA G AAC TGG CTC GG			416-441			
Fungal species: Candida glabrata							
2110 ^b	5'- <u>GCG GGC</u> CCT TAA CGA TTT C TGG ATT CA <u>G CCC GC</u>	AG CGA ATC	417	307-335			
2111	5'-GCG GGC ATG TTG AAG CCA C	CCA CCA ACG	417	419-447			
Fungal s	oecies: Candida k	rusei					
2112 ^b	5'- <u>GCG GGC</u> TTG ATG AAG TTT GGO TGA CAA TT <u>G CCC</u> GC	TTT CCT	422	318-347			
2113	5'- <u>GCG GGC</u> ACA AGG GTT GGA CT CCA AGG CA <u>G CCC GC</u>	A AGG AAA	422	419-447			
2114	5'- <u>GCG GGC</u> ATC GAT GCT ATT GA GTC AGA CC <u>G CCC GC</u>	A CCA CCT	422	505-533			

a Underlined nucleotides indicate the molecular beacon's stem.

b Sequence from databases.

c These sequences were aligned to derive the corresponding primer.

d The nucleotide positions refer to the P. aeruginosa tuf sequence fragment (SEQ ID NO. 153).

Annex XLVII: Molecular beacon internal hybridization prob s for specific d tection of tuf sequences (continued).

		Originating DNA	fragment
SEQ ID NO.	Nucleotide sequence ^a	SEQ ID Nuc	electide osition
Fungal sp	oecies: Candida lusitaniae	·	
2115 ^b	5'-GCG GGC GGT AAG TCC ACC GGT AAG ACC TTG TTG GCC CGC	424	304-330
2116	5'- <u>GCG GGC</u> GTA AGT CAC CGG TAA GAC CTT GTT G <u>GC CCG</u> <u>C</u>	424	476-502
2117	5'-GCG GGC GAC GCC ATT GAG CCA CCT TCG AGA GCC CGC	424	512-535
Fungal s	pecies: Candida parapsilosis		
2118 ^b	5'-GCG GGC TCC TTG ACA ATT TCT TCG TAT CTG TTC TTG GCC CGC	426	301-330
Fungal s	pecies: Candida tropicalis		
2119	5'-GCG GGC TTA CAA CCC TAA GGC TGT TCC ATT CGT TGC CCG C	429	357-384
2120	5'-GCG GGC AGA AAC CAA GGC TGG TAA GGT TAC CGG AGC CCG C	429	459-487
Fungal s	pecies: Cryptococcus neoform	nans	
2106	5'-GCG AGC AGA GCA CGC CCT CCT CGC CGC. TCG C		226-244 ^d
2107	5'-GCG AGC TCC CCA TCT CTG GTT GGC ACG CTC GC	623,1985,1986 ^C	390-408 ^d
Bacteria	al genus: Legionella sp.		
2083	5'-CCG CCG ATG TTC CGT AAA TTA CTT GAI GAA GGT CGA GCC GGC GG	111-112 ^a ,	488-519 ^e

a Underlined nucleotides indicate the molecular beacon's stem.

b This sequence is from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

C These sequences were aligned to derive the corresponding primer.

d The nucleotide positions refer to the C. neoformans tuf (EF-1) sequence fragment (SEQ ID NO. 623).

 $^{^{}m e}$ The nucleotide positions refer to the L. pneumophila tuf (EF-1) sequence fragment (SEQ ID NO. 112).

Annex XLVII: Molecular beacon internal hybridization probes for specific detection of tuf sequences (continued).

			Originating DNA	fragment
SEQ ID NO.	Nucleotide	sequence ^a		cleotide osition
Fungal ge	enus:	Candida sp.		
2108		AAC TTC RTC AAG AAG GTT GGT CCG CCC GC	414,417, 422,424, 426,429,624 ^b	52-80 ^c
2109	5'- <u>GCG GGC</u> GAC AA <u>G</u>	CCA ATC TCT GGT TGG AAY GGT	Same as SEQ ID NO. 2108	100-125 ^C
Bacterial group: Pseudomonads				
2121	5'- <u>CGA CCG</u> GTC G	CIA GCC GCA CAC CAA GTT CCG	153-155, 205,880,2137 ^d , 2138 ^d ,b	598-616 ^e

a Underlined nucleotides indicate the molecular beacon's stem.

b These sequences were aligned to derive the corresponding primer.

C The nucleotide positions refer to the C. albicans tuf (EF-1) sequence fragment (SEQ ID NO. 624).

d Sequence from databases.

 $^{^{}m e}$ The nucleotide positions refer to the $\it{P. aeruginosa}$ tuf sequence fragment (SEQ ID NO. 153).

Annex XLVIII: Molecular beacon internal hybridization prob s for specific detection of ddl and mtl gene sequences.

			DVI functions
		Originating	DNA fragment
SEQ ID NO. Nucleotic	de sequence ^a	SEQ ID NO.	Nucleotide position
Bacterial species	: E. faecium (ddl)		
Bacterrar species			
1237 5'- <u>GCG A(</u> TTA GO	GC CGC GAA ATC GAA GTT GCT GTA GG CTC GC	1242 ^b	334~359
Bacterial species	E. faecalis (mtl)		•
	GC GGC GTT AAT TTT GGC ACC GAA AG CTC GC	1243 ^b	631-656

a Underlined nucleotides indicate the molecular beacon's stem.

b Sequence from databases.

Annex XLIX: Internal hybridization probe for specific detection of S. aur us sequenc s of unknown coding potential.

		Originating	g DNA fragment
SEQ ID NO. Nucleotide	sequence	SEQ ID NO.	Nucleotide position
Bacterial species:	Staphylococcus aureus	;	
1234 5'-ACT AAA	TAA ACG CTC ATT CG	1244	35-54

Specific and ubiquitous primers for nucleic acid Annex L: (antimicrobial ag nts resistance genes amplification sequences).

sec	ruences).			
			Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence		SEQ ID NO.	Nucleotide position
Resistance g	ene: aac(2')-Ia	·	
1344	5'-AGC AGC AAC GAT		1348 ^a	163-186
1345 ^b	5'-CCC GCC GAG CAT	TTC AAC TAT TG	· 1348 ^a	392-414
1346	5'-GAT GTT ACG CAG	CAG GGC AGT C	1348 ^a	172-193
1347 ^b	5'-ACC AAG CAG GTT	CGC AGT CAA GTA	1348 ^a	467-490
<u>Resistance g</u>	ene: aac(3')-Ib		
1349	5'-CAG CCG ACC AAT	GAG TAT CTT GCC	1351 ^a	178-201
1350b	5'-TAA TCA GGG CAG		1351 ^a	356-379
Resistance o	ene: aac(3'	')-IIb		
1352	5'-CCA CGC TGA CAG	AGC CGC ACC G	1356 ^a	383-404
1352 1353 ^b	5'-GGC CAG CTC CCA		1356 ^a	585-606
3.254	5'-CAC GCT GAC AGA	CCC CCA CCG	1356 ^a	384-404
1354 1355 ^b	5'-ATG CCG TTG CTG		1356 ^a	606-629
Resistance o	ene: aac(3	')-IVa		
4.5.5	5'-GCC CAT CCA TTT	ככר שייי כר	1361 ^a	295~314
1357 1358 ^b	5'-GCG TAC CAA CTT		1361 ^a	517-540
	5'-TGC CCC TGC CAC		1361 ^a	356-374
1359 1360 ^b	5'-CGT ACC AAC TTG		1361 ^a	516-539
Resistance o	40	′)-VIa		
1,002			_	
1362	5'-CGC CGC CAT CGC C		1366 ^a 1366 ^a	285-306 551-574
1363 ^b	5'-CGG CAT AAT GGA G	CG CGG TGA CTG	1300-	221-274
1364	5'-TTT CTC GCC CAC G	CA GGA AAA ATC	1366 ^a	502-525
1365 ^b	5'-CAT CCT CGA CGA A	TA TGC CGC G	1366 ^a	681-702
Resistance de	ne: aac(6')	-Ia		
1367	5'-CAA ATA TAC TAA C	AG AAG CGT TCA	1371 ^a	
1368 ^b	5'-AGG ATC TTG CCA A	ATA CCT TTA T	1371 ^a	269-290
1379	5'-AAA CCT TTG TTT C	GG TCT GCT AAT	1371 ^a	153-176
	5'-AAG CGA TTC CAA		1371 ^a	320-343

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex L: Specific and ubiquitous primers for nucleic acid amplification (antimicrobial agents resistance genes sequences) (continued).

Resistance gene: 1372 SEQ ID NO. Nucleotide sequence aac(6')-Ic 5'-GCT TTC GTT GCC GAG GTC	SEQ ID NO.	Nucleotide position
1 (1) 3 -GC1 11C G11 GCC 411 GCC GMG G10	1376 ^a	
1373b 5'-CAC CCC TGT TGC TTC GCC CAC TC	1376 ^a	304-326
1374 5'-AGA TAT TGG CTT CGC CGC ACC ACA	1376 ^a	104-127
1375b 5'-CCC TGT TGC TTC GCC CAC TCC TG	1376 ^a	301-323
Resistance gene: ant(3')-Ia		
1377 5'-GCC GTG GGT CGA TGT TTG ATG TTA	1381 ^a	100-123
1377 5'-GCC GTG GGT CGA TGT TTG ATG TTA 1378b 5'-GCT CGA TGA CGC CAA CTA CCT CTG	1381 ^a	
·	1381 ^a	127-150
1379 5'-AGC AGC AAC GAT GTT ACG CAG CAG 1380b 5'-CGC TCG ATG ACG CCA ACT ACC TCT	1381 ^a	222-245
Resistance gene: ant(4')-Ia		
1382 5'-TAG ATA TGA TAG GCG GTA AAA AGC	1386 ^a	149-172
1383b 5'-CCC AAA TTC GAG TAA GAG GTA TT	1386 ^a	386-408
1384 5'-GAT ATG ATA GGC GGT AAA AAG C	1386 ^a	151-172
1384 5'-GAT ATG ATA GGC GGT AAA AAG C 1385b 5'-TCC CAA ATT CGA GTA AGA GGT A	1386 ^a	388-409
Resistance gene: aph(3')-Ia		
1387 5'-TTA TGC CTC TTC CGA CCA TCA AGC	1391 ^a	233-256
1338b 5'-TAC GCT CGT CAT CAA AAT CAC TCG	1391 ^a	488-511
1389 5'-GAA TAA CGG TTT GGT TGA TGC GAG	1391 ^a	468-491
1390b 5'-ATG GCA AGA TCC TGG TAT CGG TCT	1391 ^a	669-692
Resistance gene: aph(3')-IIa		
1392 5'-TGG GTG GAG AGG CTA TTC GGC TAT	1396 ^a	43-66
1393b 5'-CAG TCC CTT CCC GCT TCA GTG AC	1396 ^a	250-272
1394 5'-GAC GTT GTC ACT GAA GCG GGA AGG	1396 ^a	244-267
1395b 5'-CTT GGT GGT CGA ATG GGC AGG TAG	1396 ^a	386-409

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Specific and ubiquitous primers for nucleic acid Annex L: amplification (antimicrobial agents resistance genes sequences) (continued).

		Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
Resistance (gene: aph(3')-IIIa		
1397	5'-GTG GGA GAA AAT GAA AAC CTA T	1401 ^a	103-124
1397 1398 ^b	5'-ATG GAG TGA AAG AGC CTG AT	1401 ^a	355-374
1200	5'-ACC TAT GAT GTG GAA CGG GAA AAG	1401 ^a	160-183
1399 1400 ^b	5'-CGA TGG AGT GAA AGA GCC TGA TG	1401 ^a	354-376
	gene: aph(3')-VIa		
<u>Resistance</u>			40 41
1402	5'-TAT TCA ACA ATT TAT CGG AAA CAG	1406 ^a	18-41 175-197
1403 ^b	5'-TCA GAG AGC CAA CTC AAC ATT TT	1406 ^a	1/5-19/
1404	5'-AAA CAG CGT TTT AGA GCC AAA TAA	1406 ^a	36-59
1405 ^b	5'-TTC TCA GAG AGC CAA CTC AAC ATT	1406 ^a	177-200
Resistance	gene: blaCARB		
10010,001		1411 ^a	351-374
1407	5'-CCC TGT AAT AGA AAA GCA AGT AGG	1411 ^a	556-577
1408 ^b	5'-TTG TCG TAT CCC TCA AAT CAC C		
1409	5'-TGG GAT TAC AAT GGC AAT CAG CG	1411 ^a	205-227
1410 ^b	5'-GGG GAA TAG GTC ACA AGA TCT GCT T	1411 ^a	329-353
Resistance	gene: blaCMY-2		
1.110	5'-GAG AAA ACG CTC CAG CAG GGC	1416 ^a	793-813
1412 1413 ^b	5'-CAT GAG GCT TTC ACT GCG GGG	1416 ^a	975-995
	5'-TAT CGT TAA TCG CAC CAT CAC	1416 ^a	90-110
1414	5'-ATG CAG TAA TGG CAC CAT CAC	1416 ^a	439-459
_ 1415 ^b			
<u>Resistance c</u>	genes: blaCTX-M-1, blaCTX-M-2		
1417	5'-TGG TTA ACT AYA ATC CSA TTG CGG A	1423 ^a	314-338
1418 ^b	5'-ATG CTT TAC CCA GCG TCA GAT T	1423 ^a	583-604
Resistance (gene: blaCTX-M-1		
- 44.0	5'-CGA TGA ATA AGC TGA TTT CTC ACG	1423 ^a ~	410-433
1419 1420 ^b	5'-TGC TTT ACC CAG CGT CAG ATT ACG	1423 ^a	580-603
1420	-	1423 ^a	116-139
1421	5'-AAT TAG AGC GGC AGT CGG GAG GAA	1423 ^a	405-428
1422 ^b	5'-GAA ATC AGC TTA TTC ATC GCC ACG	1460	

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Specific and ubiquitous primers for nucl ic acid amplification (antimicrobial agents resistance genes Annex L: sequences) (continued).

		Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
Resistance	gene: blaCTX-M-2		
1424	5'-GTT AAC GGT GAT GGC GAC GCT AC	1428 ^a	30-52
1425 ^b	5'-GAA TTA TCG GCG GTG TTA ATC AGC	1428 ^a	153-176
1426	5'-CAC GCT CAA TAC CGC CAT TCC A	1428 ^a	510-531
1427b	5'-TTA TCG CCC ACT ACC CAT GAT TTC	1428 ^a	687-710
Resistance	gene: blaIMP		
	5'-TTT ACG GCT AAA GAT ACT GAA AAG T	1433 ^a	205-229
1429 1430 ^b	5'-TTT ACG GCT AAA GAT ACT GAA AAG T 5'-GTT TAA TAA AAC AAC CAC CGA ATA AT	1433 ^a	513-538
1430~		2	
1431	5'-TAA TTG ACA CTC CAT TTA CGG CTA A	1433 ^a	
1432 ^b	5'-ACC GAA TAA TAT TTT CCT TTC AGG CA	1433 ^a	497-522
Resistance	gene: blaOXA2		
1434	5'-CAC AAT CAA GAC CAA GAT TTG CGA T	1438 ^a	319-343
1435 ^b	5'-GAA AGG GCA GCT CGT TAC GAT AGA G	1438 ^a	532-556
Resistance	gene: blaOXA10		
4.47.6	5'-CAG CAT CAA CAT TTA AGA TCC CCA	1439 ^a	194-217
1436 1437 ^b	5'-CTC CAC TTG ATT AAC TGC GGA AAT TC	1439 ^a	479-504
Resistance		•	
		1442 ^a	281-306
1440	5'-AGA CCG TTA TCG TAA ACA GGG CTA AG 5'-TTT TTT GCT CAA ACT TTT TCA GGA TC	1442 ^a	
1441 ^b	57-TTT TIT GCT CAA ACT TIT TCA GGA TC		
Resistance (gene: blaPER-2		
1443	5'-CTT CTG CTC TGC TGA TGC TTG GC	1445 ^a	32-54
1444b	5'-GGC GAC CAG GTA TTT TGT AAT ACT GC	1445 ^a	304-329
Resistance	genes: blaPER-1, blaPER-2		
	5'-GGC CTG YGA TTT GTT ATT TGA ACT GGT	1442 ^a	414-440
1446 1447 ^b	5'-GGC CTG YGA TIT GIT AIT IGA ACT GGT 5'-CGC TST GGT CCT GTG GTG GTT TC	1442 ^a	652-674
144/		44408	522 557
1448	5'-GAT CAG GTG CAR TAT CAA AAC TGG AC	1442 ^a	
1449 ^b	5'-AGC WGG TAA CAA YCC TTT TAA CCG CT	1442 ^a	011-020

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Specific and ubiquitous primers for nucleic amplification (antimicrobial agents resistance acid Annex L: sequences) (continued).

DlaSHV TTG AGC AAA TTA AAC TA	SEQ ID NO.	Nucleotide position
TTG AGC AAA TTA AAC TA CGC AGA TAA ATC ACC AC		
CGC AGA TAA ATC ACC AC		
CGC AGA TAA ATC ACC AC		71-93
	1900ª	763-785
AAA CAC CTT GCC GAC	1900 ^a	313-333
	1900 ^a	763-785
blaTEM		
TOO COT TITT TOO GG	1927 ^a	27-46
	1927 ^a	817-838
	1927a	148-170
	1927 ^a	817-838
	21478	363-384
TGT GGC GTG TTA CGG T	2147- 2147a	484-506
catII		
TAA ATC ATC AGC GGA TA	2150 ^a	151-173
catIII		
CAG CAT TAC CTT GGG TT	2153 ^a	
ACT CTT GTA GCC GAT TA	2153 ^a	603-625
catP		
CA CAC TTT AGG AC	2156 ^a	178-197
	2156 ^a	339-361
እን መመር አጥን ምጥን ርጥር ጥጥጥ ጥኔጥ	2159a	89-115
	2159 ^a	201-221
	21.62a	48-70
	2162 ^a	231-253
	AAA CAC CTT GCC GAC GGC AGA TAA ATC ACC AC BIATEM TCC CTT TTT TGC GG TCT CAG CGA TCT GTC T GGT AAG ATC CTT GAG AG TCT CAG CGA TCT GTC T CATI TGT GGC GTG TTA CGG T AAA TGC GGA TCC ATA TT CATII CAG CAT TAC AGC C TAA ATC ATC AGC GAT TA CATII CAG CAT TAC CTT GGG TT ACT CTT GTA GCC GAT TA CATI CAG CAT TAC CTT GGG TT ACT CTT GTA GCC GAT TA CAT CAT TTT AGG AC CC GTT GCG TAT CAC TT CAC CAC CAC CAC CAC CA	### CAC AC

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Specific and ubiquitous primers for nucleic acid amplification (antimicrobial agents resistance genes sequences) (continued). Annex L:

		Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
Resistance	gene: dfrA	•	
1450	5'-ACC ACT GGG AAT ACA CTT GTA ATG GC	1452 ^a	106-131
1451b	5'-ATC TAC CTG GTC AAT CAT TGC TTC GT	1452 ^a	296-321
<u>Resistance</u>	gene: dhfrIa		
1457	5'-CAA AGG TGA ACA GCT CCT GTT T	1461 ^a	75-96
1457 1458b	5'-TCC GTT ATT TTC TTT AGG TTG GTT AA	A 1461 ^a	249-275
		· 1461ª	77-96
1459	5'-AAG GTG AAC AGC TCC TGT TT 5'-GAT CAC TAC GTT CTC ATT GTC A	1461 ^a	212 222
1560 ^b			
Resistance	genes: dhfrIa, dhfrXV		
2.452	5'-ATC GAA GAA TGG AGT TAT CGG RAA TG	1461 ^a	27-52
1453 1454b	5'-CCT AAA AYT RCT GGG GAT TTC WGG A	1461 ^a	384-408
		1461 ^a	290-313
1455	5'-CAG GTG GTG GGG AGA TAT ACA AAA 5'-TAT GTT AGA SRC GAA GTC TTG GKT AA		
1456 ^b			
<u>Resistance</u>	gene: dhfrIb		
1466	5'-AAG CAT TGA CCT ACA ATC AGT GT	1470 ^a	98-120
1467b	5'-AAT ACA ACT ACA TTG TCA TCA TTT G	AT 1470 ^a	204-230
	5'-CGT TAC CCG CTC AGG TTG GAC ATC A		183-208
1468	5'-CAT CCC CCT CTG GCT CGA TGT CG	1470 ^a	354-376
1469 ^b			
Resistance	gene: dhfrV		
	5'-GAT AAT GAC AAC GTA ATA GTA TTC CC	1475ª ^	208-233
1471 1472 ^b	5'-GCT CAA TAT CAA TCG TCG ATA TA	1475 ^a	342-364
14/2-		1 4253	95-120
1473	5'-TTA AAG CCT TGA CGT ACA ACC AGT GG	1475 ^a 1475 ^a	300-325
1474 ^b	5'-TGG GCA ATG TTT CTC TGT AAA TCT CC	14,7	300 300
Resistance	genes: dhfrIb, dhfrV		
1462	5'-GCA CTC CCY AAT AGG AAA TAC GC	1470 ^a	157-179
1462 1463b	5'-AGT GTT GCT CAA AAA CAA CTT CG	1470 ^a	405-427
		1470 ^a	139-161
1464	5'-ACG TTY GAA TCT ATG GGM GCA CT 5'-GTC GAT AAG TGG AGC GTA GAG GC	1470 ^a	328-350
1465 ^b	2 -GIC GWI WWG 100 WOC GIV ONG GO		

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Specific and ubiquitous primers for nucleic amplification (antimicrobial agents r sistanc acid Annex L: genes sequences) (continued).

		Originating	DNA fragmen
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
Resistance	gene: dhfrVI		
1476	5'-GGC GAG CAG CTC CTA TTC AAA G	1480 ^a	79-100
1477b	5'-TAG GTA AGC TAA TGC CGA TTC AAC A	1480 ^a	237-261
1478	5'-GAG AAT GGA GTA ATT GGC TCT GGA TT	1480 ^a	31-56
1478 1479 ^b	5'-GCG AAA TAC ACA ACA TCA GGG TCA T	1480 ^a	209-233
Resistance	gene: dhfrVII		
1.405	5'-AAA ATG GCG TAA TCG GTA ATG GC	1489 ^a	32-54
1485 1486 ^b	5'-CAT TTG AGC TTG AAA TTC CTT TCC TC	1489 ^a	189-214
			166-191
1487 1488 ^b	5'-AAT CGA AAA TAT GCA GTA GTG TCG AG 5'-AGA CTA TTG TAG ATT TGA CCG CCA	1489 ^a	294-317
Resistance	genes: dhfrVII, dhfrXVII		
1481	5'-RTT ACA GAT CAT KTA TAT GTC TCT	1489 ^a	
1482 ^b	5'-TAA TTT ATA TTA GAC AWA AAA AAC TG	1489 ^a	421-446
1483	5'-CAR YGT CAG AAA ATG GCG TAA TC	1489 ^a	
1484 ^b	5'-TKC AAA GCR WTT TCT ATT GAA GGA AA	1489 ^a	229-254
Resistance	gene: dhfrVIII		
1490	5'-GAC CTA TGA GAG CTT GCC CGT CAA A	1494 ^a	144-168
1491 ^b	5'-TCG CCT TCG TAC AGT CGC TTA ACA AM	4 1494 ^a	376-401
1492	5'-CAT TTT AGC TGC CAC CGC CAA TGG TT	r 1494 ^a	18-43
1493 ^D	5'-GCG TCG CTG ACG TTG TTC ACG AAG A	1494ª -	245-269
esistance o	gene: dhfrIX		
1.405	5'-TCT CTA AAC ATG ATT GTC GCT GTC	1499 ^a	7-30
1495 1496 ^b	5'-CAG TGA GGC AAA AGT TTT TCT ACC	1499 ^a	133-156
	5'-CGG ACG ACT TCA TGT GGT AGT CAG T	1499 ^a	171-195
1497 1498b	5'-TTT GTT TTC AGT AAT GGT CGG GAC CT	1499 ^a	446-471

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex L: Specific and ubiquitous primers for nucleic acid amplification (antimicrobial agents resistance genes sequences) (continued).

								Originating	DNA fragment
SEQ ID NO.	Nucleotide	sequenc	е					SEQ ID NO.	Nucleotide position
Resistance o	ene:	dhfrX	II						
1500	5'-ATC GGG	TTA TTG	GCA	ATG	GTC	СТА		1504 ^a	50-73
1501 ^b	5'-GCG GTA							. 1504 ^a	201-225
	5'-GCG GGG	, cca c cm	CAC	አጥል	ጥልሮ	10.		1504ª	304-325
1502 ₁₅₀₃ b	5'-AAC GG/	. GGA GC1	GTA	CGG	AAT	TAC	AG	1504ª	452-477
1505	3 ,2.0 00.								
Resistance of	gene:	dhfrX	III						
1505	5'-ATT TT	ר רכר אכם	СТС	ACC	GAG	AGC		1507 ^a	106-129
1505 1506 ^b	5'-CGG AT							1507 ^a	413-439
Resistance of	gene:	dhfrX	V						
1508	5'-AGA AT	G TAT TGG	TAT	TTC	CAT	СТА	TCG	1512ª	215-241
1509b	5'-CAA TG							1512 ^a	336-361
				C N N	~~~	m		1512 ^a	67-88
1510 1511 ^b	5'-TGG AG 5'-CAG AC						TCG		266-292
15112	5'-CAG AC	A CAA ICA	CAI	GAI	CCG		. 100		
<u>Resistance</u>	gene:	dhfrl	CVII						
4543	5'-TTC AA	כ כתכ אא	א ייי∨יז	א א א	CGT	CC		1517ª	201-223
1513 1514 ^b	5'-TTC AA 5'-GAA AT						т	1517 ^a	381-405
1314"								- 2	
1515	5'-GTG GT							1517 ^a	66-88 232-257
1516 ^b	5'-TCT TT	C AAA GC	A TTT	TCT	ATT	GAA	GG	1517 ^a	232-237
Resistance	gene.	mhn							
2102	5'-CAC CTT	ሮልሮ ሮሮሞ	האר ר	GA C	:G			2105 ^a	822-841
2102 2103 ^b	5'-CGA ACC					rC		2105 ^a	948-970
<u>Resistance g</u>	enes:	ereA,	ereA	2					
1528	5'-AAC TTG	AGC GAT	TTT C	GG I	ATA (cc :	rg	1530 ^a	80-105

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex L: Specific and ubiquitous primers for nuclic amplification (antimicrobial agents resistance acid sequences) (continued).

		Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
Resistance	gene: ereB		
1531	5'-TCT TTT TGT TAC GAC ATA CGC TTT T	1535 ^a	152-176
1532 ^b	5'-AGT GCT TCT TTA TCC GCT GTT CTA	1535 ^a	456-479
1533	5'-CAG CGG ATA AAG AAG CAC TAC ACA TT	1535 ^a	461-486
1534 ^b	5'-CCT CCT GAA ATA AAG CCC GAC AT	1535 ^a	727-749
Resistance	gene: gyrA		
	5'-GAA CAA GGT ATG ACA CCG GAT AAA T	1299 ^a	163-188
1340 1341 ^b	5'-GAT AAC TGA AAT CCT GAG CCA TAC G	1299ª	274-299
1341		1954 ^a	205-219
1936	5'-TAC CAC CCG CAC GGC	1954ª	309-325
1937 ^b	5'-CGG AGT CGC CGT CGA TG		
1942	5'-GAC TGG AAC AAA GCC TAT AAA AAA TCA	. 1954 ^a	148-174
1937 ^b	5'-CGG AGT CGC CGT CGA TG	1954 ^a	309-325
0040	5'-TGT GAC CCC AGA CAA ACC C	2054 ^a	33-51
2040 2041 ^b	5'-GTT GAG CGG CAG CAC TAT CT	2054 ^a	207-226
Resistance	gene:		
2098	5'-CTG AGT CAC ACC GAC AAA CGT C	2101 ^a	
2099 ^b	5'-CCA GGA CTG AAC GGG ATA CGA A	2101 ^a	1074-1095
Resistance	genes: linA, linA'		
4 = 2 =	5 / 303 TOT 307 TO 307 AVA AVA A	1540	99-123
1536 1537 ^b	5'-AGA TGT ATT AAU TGG AAA AUA AUA A 5'-CTT TGT AAT TAG TTT CTG AAA ACC A	1540 ^a	352-376
153/~			107 214
1538	5'-TTA GAA GAT ATA GGA TAC AAA ATA GAA	G 1540 ^a 1540 ^a	187-214 404-425
1539b	5'-GAA TGA AAA AGA AGT TGA GCT T	1340	404 423
Resistance	gene: linB		
1541	5'-TGA TAA TCT TAT ACG TGG GGA ATT T	1545 ^a	246-270
1541 1542 ^b	5'-ATA ATT TTC TAA TTG CCC TGT TTC AT	1545 ^a	359-384
	5'-GGG CAA TTA GAA AAT TAT TTA TCA GA	1545ª	367-392
1543 ₁₅₄₄ b	5'-GGG CAA TTA GAA AAT TAT TIA ICA GA 5'-TTT TAC TCA TGT TTA GCC AAT TAT CA	1545 ^a	579-604
15442	5 -111 IAC ICA IGI IIA GCC III III		

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex L: Specific and ubiquitous primers for nucleic acid amplification (antimicrobial agents resistance genes sequences) (continued).

	•	•	
		Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
Resistance	gene: mefA		
1546	5'-CAA GAA GGA ATG GCT GTA CTA C	1548 ^a	
1547 ^b	5'-TAA TTC CCA AAT AAC CCT AAT AAT AG	A 1548 ^a	816-842
Resistance	gene: mefE		
1549	5'-GCT TAT TAT TAG GAA GAT TAG GGG GC	1551 ^a	815-840
1550b	5'-TAG CAA GTG ACA TGA TAC TTC CGA	1551 ^a	1052-1075
Resistance	genes: mefA, mefE		
1553	5'-GGC AAG CAG TAT CAT TAA TCA CTA	1548 ^a	50-73
1552 1553 ^b	5'-CAA TGC TAC GGA TAA ACA ATA CTA TO	1548 ^a	318-343
	THE PART AND ADD THE TAC CA	AC 1548 ^a	1010-1035
1554 ₁₅₅₅ b	5'-AGA AAA TTA AGC CTG AAT ATT TAG GA 5'-TAG TAA AAA CCA ATG ATT TAC ACC G	1548 ^a	-
1555~			
Resistance	genes: mphA, mphK		
1556	5'-ACT GTA CGC ACT TGC AGC CCG ACA T	1560 ^a	
1557b	5'-GAA CGG CAG GCG ATT CTT GAG CAT	1560 ^a	214-237
1558	5'-GTG GTG GTG CAT GGC GAT CTC T	1560 ^a	583-604
1558 1559 ^b	5'-GCC GCA GCG AGG TAC TCT TCG TTA	1560 ^a	855-878
Resistance	_		
0140	5'-GCC TTA ATT TCG GAT AGT GC	2144 ^a	1831-185
2142 2143 ^D		· 2144ª	2002-2026
Resistance o	gene: parC		
1342	5'-GAT GTT ATT GGT CAA TAT CAT CCA	1321 ^a	205-229
1343 ^b	5'-AAG AAA CTG TCT CTT TAT TAA TAT CAC	GT 1321 ^a	396-425
1934	5'-GAA CGC CAG CGC GAA ATT CAA AAA G	1781	67-91
1935b	5'-AGC TCG GCA TAC TTC GAC AGG	1781	277-297
0044	5'-ACC GTA AGT CGG CCA AGT CA	2055 ^a	176-195
2044 2045 ^b	5'-GTT CTT TCT CCG TAT CGT C	2055 ^a	436-454

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex L: Specific and ubiquitous primers for nucleic acid amplification (antimicrobial agents resistance genes sequences) (continued).

		Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
Resistance	gene: ppflo-like		
2163	5'-ACC TTC ATC CTA CCG ATG TGG GTT	2165 ^a	922-945
2164 ^b	5'-CAA CGA CAC CAG CAC TGC CAT TG	2165 ^a	1136-1158
Resistance	gene: rpoB		
2065	5'-CCA GGA CGT GGA GGC GAT CAC A	2072 ^a	
206 ⁶ b	5'-CAC CGA CAG CGA GCC GAT CAG A	2072 ^a	1485-1506
Resistance	gene: satG		
1581	5'-AAT TGG GGA CTA CAC CTA TTA TGA	TG 1585 ^a	93-118
1582 ^b	5'-GGC AAA TCA GTC AGT TCA GGA GT	1585 ^a	310-332
1583	5'-CGA TTG GCA ACA ATA CAC TCC TG	1585 ^a	294-316
1584 ^b	5'-TCA CCT ATT TTT ACG CCT GGT AGG	AC 1585 ^a	388-413
Resistance	gene: sulII		
1961	5'-GCT CAA GGC AGA TGG CAT TCC C	1965 ^a	222-243
1962 ^b	5'-GGA CAA GGC GGT TGC GTT TGA T	1965 ^a	496-517
1963	5'-CAT TCC CGT CTC GCT CGA CAG T	1965 ^a	237-258
1964b	5'-ATC TGC CTG CCC GTC TTG C	1965 ^a	393-411
Resistance	e gene: tetB		
1966	5'-CAT GCC AGT CTT GCC AAC G	1970 ^a	66-84
1967 ^b	5'-CAG CAA TAA GTA ATC CAG CGA TG	1970 ^a	242-264
1050	5'-GGA GAG ATT TCA CCG CAT AG	1970 ^a	457-476
1968 1969 ^b	5'-AGC CAA CCA TCA TGC TAT TCC A	1970 ^a	721-742
Resistance	gene: tetM		
1586	5'-ATT CCC ACA ATC TTT TTT ATC AAT	AA 1590 ^a	361-386
1585 1587 ^b	5'-CAT TGT TCA GAT TCG GTA AAG TTC	1590 ^a	501-524
	5'-GTT TTT GAA GTT AAA TAG TGT TCT	т 1590 ^а	957-981
1588 1589 ^b	5'-CTT CCA TTT GTA CTT TCC CTA	1590 ^a	1172-1192

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex L: Specific and ubiquitous primers for nucleic acid amplification (antimicrobial agents resistance genes s quences) (continued).

		Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
Resistance ge	ne: vatB		
1609	5'-GCC CTG ATC CAA ATA GCA TAT A	1613 ^a	11-32
1610 ^b	5'-CCT GGC ATA ACA GTA ACA TTC TG	1613 ^a	379-401
	TO THE COLUMN TO	1613 ^a	301-322
1611	5'-TGG GAA AAA GCA ACT CCA TCT C 5'-ACA ACT GAA TTC GCA GCA ACA AT	1613 ^a	424-446
1612 ^b	5 -ACA ACT GAA TTC GCA GCA TION TO		
Resistance ge	ne: vatC		
1614	5'-CCA ATC CAG AAG AAA TAT ACC C	1618 ^a	26-47
1615 ^b	5'-ATT AGT TTA TCC CCA ATC AAT TCA	1618 ^a	177-200
2020	·	1618 ^a	241-266
1616	5'-ATA ATG AAT GGG GCT AAT CAT CGT AT 5'-GCC AAC AAC TGA ATA AGG ATC AAC	1618 ^a	463-486
1617 ^b	5'-GCC AAC AAC IGA AIA AGG AIC AAC		
Resistance ge	ene: vga		
1619	5'-AAG GCA AAA TAA AAG GAG CAA AGC	1623 ^a	641-664
1620b	5'-TGT ACC CGA GAC ATC TTC ACC AC	1623 ^a	821-843
		1623 ^a	843-868
1621	5'-AAT TGA AGG ACG GGT ATT GTG GAA AG 5'-CGA TTT TGA CAG ATG GCG ATA ATG AA	1623 ^a	975-1000
1622 ^b	5 -CGA III IGA CAG AIG GCG IIII IIIG		
Resistance ge	ene: vgaB		
1624	5'-TTC TTT AAT GCT CGT AGA TGA ACC TA	1628 ^a	354-379
1624 1625 ^b	5'-TTT TCG TAT TCT TCT TGT TGC TTT C	1628 ^a	578-602
2020		1628 ^a	663-688
1626	5'-AGG AAT GAT TAA GCC CCC TTC AAA AA	1628 ^a	849-874
1627 ^b	5'-TTA CAT TGC GAC CAT GAA ATT GCT CT	1026	045 074
Resistance gen	nes: vgb, vgh		
1.600	5'-AAG GGG AAA GTT TGG ATT ACA CAA CA	1633 ^a	73-98
	5'-GAA CCA CAG GGC ATT ATC AGA ACC	1633ª	445-468
		16228	576-596
	5'-CGA CGA TGC TTT ATG GTT TGT	1633 ^a 1633 ^a	850-875
1632 ^b	5'-GTT AAT TTG CCT ATC TTG TCA CAC TC	1000	

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex L: Specific and ubiquitous primers for nucleic acid amplification (antimicrobial agents resistance genes sequences) (continued).

		·	
		Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID	Nucleotide position
Resistance (gene: vgbB		
1634	5'-TTA ACT TGT CTA TTC CCG ATT CAG G	1882 ^a	23-47
1635b	5'-GCT GTG GCA ATG GAT ATT CTG TA	1882 ^a	267-289
	5'-TTC CTA CCC CTG ATG CTA AAG TGA	1882ª	155-178
1636 1637 ^b	5'-CAA AGT GCG TTA TCC GAA CCT AA	1882 ^a	442-464
16372		2.7.7	
	Sequencing primers		
Resistance	gene: gyrA		
1290	5'-GAY TAY GCI ATG ISI GTI ATH GT	1299 ^a	70-83
1290 1292b	5'-ARI SCY TCI ARI ATR TGI GC	1299 ^a	1132-1152
		*****	100-123
1291	5'-GCI YTI CCI GAY GTI MGI GAY GG	1299 ^a 1299 ^a	1132-1152
1292 ^b	5'-ARI SCY TCI ARI ATR TGI GC	1299-	1132-1132
1293	5'-ATG GCT GAA TTA CCT CAA TC	1299 ^a	1-21
1294 ^b	5'-ATG ATT GTT GTA TAT CTT CTT CAA	1299 ^a	2626-2651
1295 ^b	5'-CAG AAA GTT TGA AGC GTT GT	1299 ^a	1255-1275
1296	5'-AAC GAT TCG TGA GTC AGA TA	1299 ^a	1188-1208
1297	5'-CGG TCA ACA TTG AGG AAG AGC T	1300 ^a	29-51
1298b	5'-ACG AAA TCG ACC GTC TCT TTT TC	1300 ^a	415-437
Resistance	gene: gyrB		
1301	5'-GTT MGT AWT MGT CCT GST ATG TA	1307ª	J_82-105
1302b	5'-TAI ADI GGI GGI KKI GCI ATR TA	1307ª	1600-1623
1302	5'-GGI GAI GAI DYI MGI GAR GG	1307 ^a	955-975
1304 ^b	5'-CIA RYT TIK YIT TIG TYT G	1307 ^a	1024-1043
1305	5'-ATG GTG ACT GCA TTG TCA GAT G	1307 ^a	1-23
1305 1306 ^b	5'-GTC TAC GGT TTT CTA CAA CGT C	1307 ^a	1858-1888
		•	

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex L: Specific and ubiquitous primers for nucleic acid amplification (antimicrobial agents resistance genes sequences) (continued).

		Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
	Sequencing primers (continued)		
Resistance	gene: parC		
1308	5'-ATG TAY GTI ATI ATG GAY MGI GC	1320 ^a	67-90
1309 ^b	5'-ATI ATY TTR TTI CCY TTI CCY TT	1320 ^a	1993-2016
1310	5'-ATI ATI TSI ATI ACY TCR TC	1320 ^a	1112-1132
1311 ^b	5'-GAR ATG AAR ATI MGI GGI GAR CA	1320 ^a	1288-1311
1312	5'-AAR TAY ATI ATI CAR GAR MGI GC	1321 ^a	67-90
1312 1313b	5'-AMI AYI CKR TGI GGI TTI TTY TT	1321 ^a	2212-2235
1314	5'-TAI GAI TTY ACI GAI SMI CAR GC	1321 ^a	1228-1251
1315 ^b	5'-ACI ATI GCI TCI GCY TGI KSY TC	1321 ^a	1240-1263
1316	5'-GTG AGT GAA ATA ATT CAA GAT T	1321 ^a	1-23
1317b	5'-CAC CAA AAT CAT CTG TAT CTA C	1321 ^a	2356-2378
0-	5'-ACC TAY TCS ATG TAC GTR ATC ATG GA	1320 ^a	58-84
1318 1319 ^b	5'-AGR TCG TCI ACC ATC GGY AGY TT	1320 ^a	832-855
1319~	3 -Mark 160 161 Mee 1110 000 Me		
<u>Resistance</u>	gene: parE		
*200	5'-RTI GAI AAY ISI GTI GAY GAR G	1328 ^a	133-155
1322 1325 ^b	5'-RTT CAT YTC ICC IAR ICC YTT	1328 ^a	1732-1752
1323	5'-ACI AWR SAI GGI GGI ACI CAY G	1328 ^a	829-850
135 4 5°p	5 [-CCITCCITGCITSWR TCITCCYTCT	175000	1280-130200
1326	5'-TGA TTC AAT ACA GGT TTT AGA G	1328 ^a	27-49
1326 1327 ^b	5'-CTA GAT TTC CTC CTC ATC AAA T	1328 ^a	1971-1993

a Sequence from databases.

b These sequences are from the complementary DNA strand of the sequence of the originating fragment given in the Sequence Listing.

Annex LI: Internal hybridization probes for specific det ction of antimicrobial agents resistance genes sequences.

		Originating I	NA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
Resistance	gene: aph3'VIa		
2252	5'-CCA CAT ACA GTG TCT CTC	1406 ^a	149-166
Resistance	gene: blaSHV		
1886	5'-GAC GCC CGC GCC ACC ACT	1900 ^a	484-501
1887	5'-GAC GCC CGC GAC ACC ACT A	1899 ^a	514-532
1888	5'-GAC GCC CGC AAC ACC ACT A	1901 ^a	514-532
1889	5'-GTT CGC AAC TGC AGC TGC TG	1899 ^a	593-612
1890	5'-TTC GCA ACG GCA GCT GCT G	1899 ^a	594-612
1891	5'-CCG GAG CTG CCG AIC GGG	1902 ^a	692-709
1892	5'-CGG AGC TGC CAA RCG GGG	1903 ^a	693-710
1893	5'-GGA GCT GGC GAR CGG GGT	1899 ^a	694-711
1894	5'-GAC CGG AGC TAG CGA RCG	1904 ^a	690-707
1895	5'-CGG AGC TAG CAA RCG GGG T	1905 ^a	693-711
1896	5'-GAA ACG GAA CTG AAT GAG GCG	1899 ^a	484-504
1896	5'-CAT TAC CAT GGG CGA TAA CAG	1899 ^a	366-386
1898	5'-CCA TTA CCA TGA GCG ATA ACAG	1899 ^a	365-386
Resistance	gene: blaTEM		
	5'-ATG ACT TGG TTA AGT ACT CAC C	1928 ^a	293-314
1909	5'-ATG ACT TGG TTG AGT ACT CAC C	1927 ^a	293-314
1910	5'-CCA TAA CCA TGG GTG ATA ACA C	1928 ^a	371-392
1911	5'-CCA TAA CCA TGG GTG ATA ACA C	1927 ^a	371-392
1912		1928 ^a	475-494
1913	5'-CGC CTT GAT CAT TGG GAA CC	1927 ^a	475-494
1914	5'-CGC CTT GAT CGT TGG GAA CC 5'-CGC CTT GAT AGT TGG GAA CC	1929 ^a	475-494
1915	5'-CGT GGG TCT TGC GGT ATC AT	1927 a -	712-731
1916	5'-CGT GGG TCT GGC GGT ATC AT	1930ª	712-731
1917	5'-GTG GGT CTC ACG GTA TCA TTG	1927 ^a	713-733
1918		1931 ^a	712-732
1919	5'-CGT GGG TCT CTC GGT ATC ATT	1927 ^a	712-731
1920	5'-CGT GGI TCT CGC GGT ATC AT 5'-CGT GGG TCT AGC GGT ATC ATT	1932 ^a	713-733
1921	5'-CGT GGG TCT AGC GGT ATC ATT 5'-GTT TTC CAA TGA TTA GCA CTT TTA	1927 ^a	188-211
1922		1927 ^a	188-211
1923	5'-GTT TTC CAA TGA TAA GCA CTT TTA 5'-GTT TTC CAA TGC TGA GCA CTT TT	1932 ^a	188-210
1924	5'-GTT TTC CAA TGC TGA GCA CTT TT 5'-CGT TTT CCA ATG ATG AGC ACT TT	1927 ^a	187-209
1925	5'-CGT TTT CCA ATG ATG AGC ACT IT 5'-GTT TTC CAA TGG TGA GCA CTT TT	1933ª	188-210
1926	5'-TGG AGC CGG TGA GCG TGG	1927 ^a	699-716
2006	2100 WC COO LOW CCO 100		

a Sequence from databases.

Annex LI: Internal hybridization probes for specific detection of antimicrobial agents resistance gens sequences (continu d).

		Originating I	NA fragment
SEQ ID NO.	Nucleotide sequence	SEQ ID NO.	Nucleotide position
Resistance	gene: blaTEM (continued)		
2007	5'-TGG AGC CAG TGA GCG TGG	2010 ^a	699-716
2007	5'-TCT GGA GCC GAT GAG CGT G	1929 ^a	697-715
2009	5'-CTG GAG CCA GTA AGC GTG G	2011 ^a	698-716
2141	5'-CAC CAG TCA CAG AAA AGC	1927 ^a	311-328
Resistance	gene: dhfrla		
2253	5'-CAT TAC CCA ACC GAA AGT A	1461 ^a	158-176
Resistance	gene: embB		
2104	5'-CTG GGC ATG GCI CGA GTC	2105 ^a	910-927
Resistance	gene: gyrA		
1222	5'-TCA TGG TGA CTT ATC TAT TTA TG	1299 ^a	240-263
1333	5'-CAT CTA TTT ATA AAG CAA TGG TA	1299 ^a	251-274
1334	5'-CTA TTT ATG GAG CAA TGG T	1299 ^a	254-273
1335	5'-GTA TCG TTG GTG ACG TAA T	1299 ^a	206-224
1940	5'-GCT GGT GGA CGG CCA G	1954 ^a	279-294
1943 1945	5'-CGG CGA CTA CGC GGT AT	1954 ^a	216-232
1945	5'-CGG CGA CTT CGC GGT AT	1954 ^a	216-232
	5'-CGG TAT ACG GCA CCA TCG T	1954 ^a	227-245
1947 1948	5'-GCG GTA TAC AAC ACC ATC G	1954 ^a	226-244
	5'-CGG TAT ACG CCA CCA TCG T	1954 ^a	227-245
1949	5'-CAC GGG GAT TTC TCT ATT TA	2054 ^a	103-122
2042 2043	5'-CAC GGG GAT TAC TCT ATT TA	2054 ^a	103-122
Resistance o		•	
2100	5'-GCG AGA CGA TAG GTT GTC	2101 ^a	1017-1034
Resistance (gene: parC		
1226	5'-TGG AGA CTA CTC AGT GT	1321 ^a	232-249
1336 1337	5'-TGG AGA CTT CTC AGT GT	1321 ^a	232-249
	5'-GTG TAC GGA GCA ATG	1321 ^a	245-260
1338	5'-CCA GCG GAA ATG CGT	1321 ^a	342-357
1339	5'-GCA ATG GTC CGT TTA AGT	1321 ^a	253-270
1941	5'-TTT CGC CGC CAT GCG TTA C	1781	247-265
1944 1950	5'-GGC GAC ATC GCC TGC	1781	.137-151
1951	5'-GGC GAC AGA GCC TGC TA	1781	137-153
1901	3 333 333 333 333		

a Sequence from databases.

Annex LI: Internal hybridization probes for sp cific detection of antimicrobial ag nts resistance genes sequences (continued).

		·		
		Originating DNA fragment		
SEQ ID NO.	Nucleotide sequence	SEQ ID Nucleotide NO. position		
Resistance	gene: parC (continued)			
1952	5'-CCT GCT ATG GAG CGA TGG T	1781 147-165		
1953	5'-CGC CTG CTA TAA AGC GAT GGT	1781 145-165		
2046	5'-ACG GGG ATT TTT CTA TCT AT	2055 ^a 227-246		
Resistance	gene: rpoB			
2067	5'-AGC TGA GCC AAT TCA TGG	2072 ^a 1304-1321		
2068	5'-ATT CAT GGA CCA GAA CAA C	2072 ^a 1314-1332		
2069	5'-CGC TGT CGG GGT TGA CCC	2072 ^a 1334-1351		
2070	5'-GTT GAC CCA CAA GCG CCG	2072 ^a 1344-1361		
2071	5'-CGA CTG TCG GCG CTG GGG	2072 ^a 1360-1377		
Resistance	gene: tetM			
2254	5'-ACC TGA ACA GAG AGA AAT G	1590 ^a 1062-1080		

a Sequence from databases.

Annex LII: Molecular beacon internal hybridization probes for specific det ction of atpD sequences.

		Originating	DNA fragment
SEQ ID NO.	Nucleotide sequence ^a	SEQ ID NO.	Nucleotide position
Bacterial	species: Bacteroides fragilis	-	
2136	5'-CCA ACG CGT CCT CAA TCA TTT CTA ACT TCT ATG GCC GGC GTT GG	г 929 .	353-382
Bacterial	species: Bordetella pertussis		
2182	5'-GCG CGC CAA CGA CTT CTA CCA CGA AAT GG. AGA GTC GCG CGC	A 1672	576-605
Bacterial	l group: Campylobacter jejuni	and C. col	i
2133	5'- <u>CCA</u> <u>CGC</u> ACA WAA ACT TGT TTT AGA AGT AGC AGC WCA <u>GCG</u> <u>TGG</u>	1576, 1600,1849, 1863,2139 ^b ,c	44-73 ^d
Fungal s	pecies: Candida glabrata		
2078	5'- <u>CCG AGC</u> CTT GGT CTT CGG CCA AAT GAA C <u>G</u> <u>TCG</u> <u>G</u>	<u>sc</u> 463	442-463
Fungal s	pecies: Candida krusei		
2075	5'-CCG AGC CAG GTT CTG AAG TCT CTG CAT TA TAG GTG CTC GG	AT 468	720-748
Fungal s	pecies: Candida lusitaniae		
2080	5'- <u>CCG AGC</u> CGA AGA GGG CCA AGA TGT C <u>GC</u> TC G	CG 470	520-538
Fungal s	pecies: Candida parapsilosis	1	
2079	5'-CCG AGC GTT CAG TTA CTT CAG TCC AAG CCG GCT CGG	472	837-860
Fungal spe	ecies: Candida tropicalis		
2077	5'-CCG AGC AAC CGA TCC AGC TCC AGC TAC GCT CGG	475	877-897
Bacterial	species: Klebsiella pneumoniae		
2281	5'-CCC CCA GCT GGG CGG CGG TAT CGA TGG GGG	317	40-59

a Underlined nucleotides indicate the molecular beacon's stem.

b Sequence from databases.

C These sequences were aligned to derive the corresponding primer.

d The nucleotide positions refer to the C. jejuni atpD sequence fragment (SEQ ID NO. 1576).

Annex LII: Molecular beacon internal hybridization probes for specific detection of atpD sequences (continued).

				Originating DNA fragment		
SEQ ID NO.	Nucleotide seque	nce ^a	SEQ ID NO.	Nucleotide position		
Fungal g	enus:	Candida sp.				
2076	5'- <u>CCG AGC</u> YGA Y RGC <u>GCT CGG</u>	AA CAT TTT CAG ATT CAC CCA	460-478, 663 ^b	697-723 ^C		

a Underlined nucleotides indicate the molecular beacon's stem.

b These sequences were aligned to derive the corresponding primer.

 $^{^{\}text{C}}$ The nucleotide positions refer to the $^{\text{C}}$. albicans atpD sequence fragment (SEQ ID NO. 460).

Annex LIII: Internal hybridization probes for sp cific detection of atpD sequences.

			Originating DN	A fragment
SEQ ID NO.	Nucleotide	sequence	SEQ ID NO.	Nucleotide position
Bacterial	species:	Acinetobacter baumanı	aii	
2169	5'-CCC GTT	TGC GAA AGG TGG	243	304-321
Bacterial	species:	Klebsiella pneumonia	9	
2167	5'-CAG CAG	CTG GGC GGC GGT	317	36-53

Annex LIV: Internal hybridization probes for specific det ction of ddl and mtl sequenc s.

							Originating	DNA fragment
SEQ ID NO. Nucleotide sequence						SEQ ID NO.	Nucleotide position	
Bacterial	species:	En	tero	ococ	cus	faecium	(đ đ1)	
2286	5'-AGT	TGC TGT	ATT	AGG	AAA	TG	2288 ^a	784-803
2287	5'-TCG	AAG TTG	CTG	TAT	TAG	GA	2288 ^a	780-799
Bacterial	species:	En	ter	0000	cus	faecali	s mt1)	
2289	5'-CAC	CGA AGA	AGA	TGA	AAA	AA	1243ª	264-283
2290	5'-TGG	CAC ÇGA	AGA	AGA	TGA		1243 ^a	261-278
2291		TTG GCA				A	1243 ^a	257-275

a Sequence from databases.